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(54) Title: LITHIUM MANGANESE OXIDE AND METHODS OF MANUFACTURE

(57) Abstract

This invention relates to metal-doped lithium manganese oxide and methods of its production. In particular, it relates to stable lithium manganese oxide in monoclinic form, doped with one or more metal ion, the preferred dopants being one or both of aluminium and chromium. The method of the invention involves the synthesis of stable metal-doped lithium manganese oxide in monoclinic form by high temperature solid state reaction.

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LITHIUM MANGANESE OXIDE AND METHODS OF MANUFACTURE

TECHNICAL FIELD

This invention relates to metal doped lithium manganese oxide and methods of its production.

BACKGROUND

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Given the huge commercial success of rechargeable lithium batteries there is considerable interest in the development of improved intercalation compounds for use as the electrodes in such batteries. Appropriate compounds must possess a sufficient capacity for providing energy, must have appropriate electric conductivity and, preferably, should be stable at elevated temperatures. Furthermore, they must be able to be produced in commercial quantities at a viable cost.

The trivalent manganese compound LiMnO₂ has been identified as a possible cathode material for lithium ion batteries. In particular, monoclinic (m-) LiMnO₂ (space group C2/m), having a layered rock salt structure similar to LiCoO₂ and LiNiO₂, exhibits promising electrochemical properties. However, LiMnO₂ obtained via conventional solid state reaction usually has an orthorhombic (o-LiMnO₂) structure (space group Pmnm), since m-LiMnO₂ is not usually stable under these synthesis conditions.

To date, m-LiMnO₂ has been obtained by an ion exchange reaction of lithium salts with NaMnO₂ as precursor, or by a mixed alkaline-hydrothermal reaction (Armstrong & Bruce, Nature <u>381</u>, 499 (1996); Tabuchi et. al., J. Electrochem. Soc, <u>145</u>, L49 (1998)).

International patent application no. PCT/US97/18839, by Massachusetts
Institute of Technology, discloses compositions having a general formula Li_xM_yN_xO₂,

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and including the compound LiAl_{0.25}Mn_{0.75}O₂, being a layered monoclinic structure. The added aluminium apparently stabilises the layered monoclinic modification and allows direct synthesis of the compound. Chiang et al., in Electrochem. Solid State Lett. 1, 13 (1998) describe the preparation of both LiAl_{0.25}Mn_{0.75}O₂ and LiAl_{0.05}Mn_{0.95}O₂ in the layered monoclinic structure. However, the synthesis route used to prepare the compounds in the above references (PCT/US97/18839 and Chiang et al.) is based on the coprecipitation of the precursor hydroxides out of aqueous nitrate solution and subsequent freeze drying of the obtained hydroxide mixture. The dry mixture is then fired at 945°C in an inert atmosphere.

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This synthesis method is complex and expensive and the nitrates of aluminium and manganese may be carcinogens and mutagens. Thus, the method may not be appropriate for commercial production.

Davidson et al. in United States patent No. 5,370,949 and J. Power Sources

54, 205 (1995) disclose compounds of composition Li₂Cr_xMn_{2-x}O₄ prepared by high temperature solid state reactions of lithium carbonate with chromium (III) oxide and manganese (IV) oxide, or of LiCrO₂ with o-LiMnO₂. Compounds for which x ≥ 1.5 apparently had a layered hexagonal structure similar to LiCrO₂. Compounds for which
 0.1 ≤ x ≤1.25 gave X-ray diffraction patterns which could be indexed to a tetragonal cell based on a structure like the lithiated spinel-related phase Li₂Mn₂O₄. Dahn et al.

in J. Electrochem. Soc. <u>145</u>, 851 (1998) showed that the data described by Davidson et al. in US 5,370,949 and J. Power Sources <u>54</u>, 205 (1995) are better explained by a

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monoclinic layered crystal structure.

The data published by Davidson et al. and Dahn et al., if taken in combination, therefore demonstrate that chromium can be used in a solid state reaction to stabilise a layered monoclinic structure for $\text{LiCr}_x \text{Mn}_{1-x} \text{O}_2$ for $x \ge 0.05$ (dividing the subscripts in the formulae given by Davidson et al. and Dahn et al. by a factor of 2 for consistency with other descriptions).

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The selection of element or elements doped into LiMnO₂ and the proportions of dopant may be important in determining the extent to which a stable LiMnO₂ composition may be produced with the required electrochemical properties by a commercially viable process.

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Moreover, a combination of dopants used to stabilise a layered monoclinic phase of LiMnO₂ may facilitate the synthesis procedure and/or provide electrochemical advantages over the prior art compounds using only aluminium or chromium as the dopant.

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OBJECTS OF THE INVENTION

It is therefore an object of the present invention to provide a method of synthesising lithium manganese oxide intercalation compounds which reduces or overcomes the abovementioned problems, or which at least provides the public with a useful alternative.

It is a further and/or alternative object of the present invention to provide lithium manganese oxide intercalation compounds having improved characteristics, or at least to provide the public with useful alternative compounds.

Other objects of the present invention may become apparent from the following description which is given by way of example only.

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SUMMARY OF THE INVENTION

According to one aspect of the present invention there is provided a metal-doped lithium manganese oxide composition having the formula $\text{LiM}^{1}_{m1}\text{M}^{2}_{m2}...\text{M}^{x}_{mx}\text{Mn}_{z}\text{O}_{2} \text{ wherein M}^{1}, \text{M}^{2}...\text{M}^{x} \text{ are each different metal ions,}$ m1+m2+...mx+z is substantially equal to 1, m1+m2+...mx is greater than zero but less $\text{than 0.75 and z and at least m}_{1} \text{ are each greater than zero, providing that where }$ $\text{m2...mx are zero M}^{1} \text{ is Cr and 0<m1<0.05 or M}^{1} \text{ is Ga.}$

Preferably, M¹, M²...M^x are each selected to have an ionic radius in their relevant oxidation state, similar to or less than that of Mn³⁺.

Preferably, M¹, M²...M^x are each selected from the group consisting of Al, Ga, Cr, Co, Fe, B, V, Ti and Ru.

Preferably, M¹ may be Al.

Preferably, M² may be selected from Cr, Co, Fe, B, V, Ti and Ru and is most preferably Cr.

Preferably, M^1 , M^2 ... M^x each has the property of existing in a minimally distorted octahedral sphere of co-ordination.

Preferably, $M^1, M^2...M^x$ each has a formal oxidation state of 3^+ .

Preferably, $m_1 + m_2 + ... m_x$ may be greater than zero but less than about 0.25.

Preferably the composition is in predominantly monoclinic form.

According to a further aspect of the invention there is provided a method of synthesising a metal-doped lithium manganese oxide composition having the formula $\text{LiM}^1_{m1} \text{ M}^2_{m2}...\text{M}^x_{mx} \text{ Mn}_z \text{ O}_2$ wherein M¹, M²...M^x are each different metal ions,

m1+m2+...mx + z is substantially equal to 1, m1+m2+...mx is greater than zero but less than 0.75 and z and at least m1 are each greater than zero, providing that where m2...mx are zero and M^1 is Cr then 0<m1<0.05, in stable form by solid state reaction.

5 Preferably, the solid state reaction includes:

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 mixing reagents including a lithium compound, a manganese compound and at least one additional metal compound, each compound selected from an oxide, a hydroxide and/or a carbonate, and

- heating the mixture in a reactor under inert atmosphere, to a temperature exceeding 800°C for more than 2 hours.

Preferably, the lithium compound may be lithium carbonate.

Preferably, the manganese compound may be manganese oxide.

Preferably, the or each metal of the at least one additional metal compound may be selected to have an ionic radius in its relevant oxidation state, similar to or less than that of Mn³⁺.

Preferably, the or each metal of the at least one additional metal compound may be selected from the group consisting of Al, Ga, Cr, Co, Fe, B, V, Ti and Ru.

Preferably, the at least one additional metal compound includes aluminium oxide. Preferably, a transitional alumina.

Preferably, the at least one additional metal compound alternatively, or in addition, includes chromium oxide.

Preferably, the method may further include ball milling the mixed reagents.

Preferably, the mixture may be heated to a temperature in the range substantially 900°C to 1080°C. More preferably substantially 1000°C to 1080°C for a period of at least 5 hours.

Preferably, the inert atmosphere may have a sufficiently low pO₂ to facilitate the formation of manganese in its 3+ oxidation state.

Preferably, the pO₂ may be substantially 10⁻⁵ atm or less.

Preferably, the mixture may be heated in a reactor having a high temperature stable lining.

According to a further aspect of the invention there is provided a method of synthesising a metal-doped lithium manganese oxide composition of the formula $\text{LiM}^1_{\text{ml}}\text{M}^2_{\text{m2}...}\text{M}^x_{\text{mx}}\text{Mn}_z\text{O}_2$ wherein M¹, M²...M^x are each different metal ions, m1+m2+...mx+z is substantially equal to 1, m1+m2+...mx is greater than zero but less than 0.75 and z and at least m1 are each greater than zero, in stable predominantly monoclinic form, the method including the steps of:

- mixing reagents including a lithium compound, a manganese compound
 and at least one additional metal compound, each compound selected
 from an oxide, a hydroxide and/or a carbonate,
 - heating the mixture in air to a temperature ≥ 200°C for at least 5 hours to produce oxygen-rich lithium manganese oxides,
 - further heating the mixture in a reactor under inert atmosphere to a temperature exceeding 800°C for at least 2 hours.

Preferably the heating in air may be at a temperature in the range 600 to 850°C. Most preferably substantially 700°C.

Preferably the heating under inert atmosphere may be at a temperature of substantially 1000 to 1080°C.

According to a further aspect of the present invention there is provided a method of synthesising a metal doped lithium manganese oxide composition substantially as herein described and with reference to the accompanying examples.

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Other aspects of the present invention may become apparent from the following description which is given by way of example only and with reference to the accompanying examples.

15 BRIEF DESCRIPTION OF FIGURES

Figure 1:

Effect of temperature on LiMnO₂ production by high temperature solid state reaction with metal doping in a

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steel reactor in accordance with Example I.

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Figure 2:

Effect of reaction time on LiMnO₂ production in

accordance with Example I.

Figures 3 A-F:

X-ray diffraction patterns showing the effects of temperature on the production and phase of LiMnO₂ from a high temperature solid state reaction with metal doping in an alumina-lined reactor in accordance with

Example II:

• - o-LiMnO₂

+- m-LiMnO₂

 $o-\ Li_2MnO_3$

* - Mn₃O₄

5	Figures 4 A-D:	X-ray diffraction patterns showing the effects of chromium doping on the production and phase of LiMnO ₂ from a high temperature solid state reaction with metal doping in an alumina-lined reactor, in accordance with Example IV:
10		 • - o-LiMnO₂ + - m-LiMnO₂ o - o-Li₂MnO₃ * - Mn₃O₄ ◆ - LiCrO₂
15	Figure 5:	X-ray diffraction patterns of $LiCr_xMn_{1-x}O_2$ for $x=0.01$, 0.03, 0.05, 0.10 and 0.20, prepared according to Example V (+ = m-LiMnO ₂ , o = o-LiMnO ₂).
20	Figure 6:	X-ray diffraction pattern of LiAl _{0.05} Mn _{0.95} O ₂ prepared according to Example VI (+ = m-LiMnO ₂ , o = o-LiMnO ₂).
	Figure 7:	Schematic representation of the cell construction as used in Examples VI, VII, VIII and X.
25	Figure 8:	X-ray diffraction pattern of LiAl _{0.10} Mn _{0.90} O ₂ prepared according to Example VII ($o = o$ -LiMnO ₂ , all other peaks m-LiMnO ₂).
30	Figures 9A-D:	X-ray diffraction patterns of (A) LiAl $_{0.05}$ Cr $_{0.01}$ Mn $_{0.94}$ O $_2$, (B) LiAl $_{0.05}$ Cr $_{0.03}$ Mn $_{0.92}$ O $_2$, (C) LiAl $_{0.05}$ Cr $_{0.05}$ Mn $_{0.90}$ O $_2$,

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and (D) LiAl $_{0.02}$ Cr $_{0.05}$ Mn $_{0.93}$ O $_2$ prepared according to Example VIII (o = o-LiMnO $_2$).

Figures 10A&B:

X-ray diffraction patterns of (A) $Li_{0.5}Cr_{0.05}Mn_{0.95}O_2$ and

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(B) LiCr_{0.05}Mn_{0.95}O₂ prepared according to Example

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Figures 11A&B:

X-ray diffraction patterns of LiAl_{0.05}Mn_{0.95}O₂ prepared

according to Example X (A) after air step (* = spinel,

 $\# = \text{Li}_2\text{MnO}_3$), (B) after nitrogen step (o = o-LiMnO₂).

Figures 12A&B:

X-ray diffraction patterns of (A) Li_{0.5}Ga_{0.10}Mn_{0.90}O₂ and

(B) LiGa_{0.10}Mn_{0.90}O₂ prepared according to Example XI

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 $(o = o-LiMnO_2).$

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Figures 13A-D:

X-ray diffraction patterns of (A)

 $Li_{0.5}Cr_{0.05}Ga_{0.05}Mn_{0.90}O_2$, (B) $LiCr_{0.05}Ga_{0.05}Mn_{0.90}O_2$, (C)

 $Li_{0.5}Cr_{0.05}B_{0.05}Mn_{0.90}O_2$, and (D) $LiCr_{0.05}B_{0.05}Mn_{0.90}O_2$

prepared according to Example XII (o = o-LiMnO₂).

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DETAILED DESCRIPTION OF THE INVENTION

Investigations were carried out to identify new metal-doped LiMnO₂

compositions and provide a robust, simple and commercially viable solid state
synthesis of aluminium, chromium and/or other metal doped layered monoclinic mLiMnO₂, which could have commercial significance as a cathode in lithium ion
batteries.

The basic reaction underlying the synthesis method and compositions of the present invention involves the conversion of, for example, lithium carbonate and

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manganese oxide to lithium manganese oxide. One possible reaction scheme would be:

 $Li_{2}CO_{3} \rightarrow Li_{2}O + CO_{2}$ $2 MnO_{2} \rightarrow Mn_{2}O_{3} + \frac{1}{2}O_{2}$ $5 Li_{2}O + Mn_{2}O_{3} \rightarrow 2LiMnO_{2}.$

The invention involves methods of synthesising stable forms of metal-doped LiMnO₂, in layered monoclinic form, by solid state reaction. The invention also relates to specific stable compositions formed by these methods.

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Preferred, stable compositions were produced with the starting reagents lithium carbonate (Li₂CO₃), manganese (IV) oxide (MnO₂) and aluminium oxide (Al₂O₃) heated in a reactor under inert atmosphere to a temperature in the range 800°C to 1080°C for a period of at least 2 hours. The formation of LiMnO₂ in monoclinic form, may be facilitated by raising the temperature to between about 950°C and 1080°C, and using a transitional alumina as the source of aluminium.

Other preferred, stable compositions were produced with the starting reagents lithium carbonate (Li₂CO₃), manganese (IV) oxide (MnO₂), aluminium oxide (Al₂O₃) and chromium oxide (Cr₂O₃) heated in a reactor under inert atmosphere to a temperature in the range 800°C to 1080°C for a period of at least 2 hours.

Other preferred, stable compositions were produced in a two-stage reaction by firstly reacting lithium carbonate (Li₂CO₃) and manganese carbonate (MnCO₃) with one or more of aluminium oxide (Al₂O₃), chromium oxide (Cr₂O₃), gallium oxide (Ga₂O₃) and boric acid (H₃BO₃) in air at a temperature of 600-850 °C to produce oxygen-rich lithium manganese oxides, followed by a second heating step under nitrogen at 800-1080 °C.

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In this two-stage reaction, if the lithium, manganese and dopant reagents in the first stage are combined in the appropriate ratios to give a product with a

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lithium/(manganese + dopant) molar ratio of 1, the reaction in air will produce a two phase mixture of a spinel phase $\text{Li}_x \text{Mn}_{2-x} \text{O}_4$ and rock salt phase $\text{Li}_2 \text{MnO}_3$. Both phases may contain the dopant elements in their crystalline structure. When this product is heated under N_2 flow, the two phases react together to give doped LiMnO_2 according to the reaction scheme:

$$LiM_{2x}Mn_{2-2x}O_4 + Li_2M_xMn_{1-x}O_3 \rightarrow 3 LiM_xMn_{1-x}O_2 + \frac{1}{2}O_2$$

where M may be one or more dopant elements.

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Alternatively, the reagents in the first stage can be combined in the appropriate ratios to give a product with a lithium/(manganese + dopant) ratio of approximately 0.5. The product of the reaction in air will then be a single-phase doped spinel oxide. This can then be combined with additional Li reactant in the second stage under N_2 to give doped LiMnO₂, e.g.:

$$LiM_{2x}Mn_{2-2x}O_4 + \frac{1}{2}Li_2CO_3 \rightarrow 2LiM_xMn_{1-x}O_2 + \frac{1}{2}CO_2 + \frac{1}{2}O_2$$

An advantage of the two-stage method is that the dopant precursor compounds are reacted with the manganese precursor compound in air, which provides more reactive conditions for the decomposition of some metal salts than an inert atmosphere such as N₂. This may allow the selection of metal salts as starting compounds which may be difficult to react together to give a single-phase product under an inert atmosphere. Spinel LiMn₂O₄ can be prepared with a large number of different dopants at a range of concentrations. Many of these doped LiMn₂O₄ compounds are known and have been well characterised in terms of the effects of the dopant element on the crystal structure. This allows for ready identification of the doped spinel intermediate phase by techniques such as X-ray powder diffraction. The relatively facile formation of doped spinel phases in air then provides an atomic-level dispersion of the dopant element in the manganese oxide lattice before proceeding to the conversion step under N₂. The conversion step under N₂ therefore involves

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essentially lithium diffusion into the spinel phase from the Li₂MnO₃ phase or additional Li reactant, some reduction of Mn⁴⁺ to Mn³⁺, and an accompanying crystal phase transformation.

It will be appreciated by those skilled in the art that different lithium, manganese and dopant compounds may also be employed as the starting reagents, but generally selected from carbonates, oxides and hydroxides.

It is considered that the metal ions most preferred for doping of the LiMnO₂ should be selected by several criteria, firstly ionic radius, and secondly appropriate charge state. The incorporation of a metal ion with an ionic radius similar to or less than that of Mn³⁺ is likely to mitigate the distortion of the octahedral geometry around the Mn³⁺. Thus, preferably, the metal ion or ions should have an ionic radius, in their relevant oxidation state, similar to or less than that of Mn³⁺. Hence, as an alternative to, or in addition to aluminium, the dopants may be selected from, for example, B, Ga, Cr, Co, Fe, V, Ti and Ru. Preferably, the metal ion or ions may also have the property of existing in a minimally distorted octahedral sphere of co-ordination.

20 Example I

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Exact stoichiometric amounts (based on LiAl_{0.05}Mn_{0.95}O₂) of lithium carbonate (99.9% pure), manganese (IV) oxide (90+% pure) and aluminium oxide (99.8% pure) were weighed out, mixed and subsequently homogenised for 30 minutes using a vibrating table. These mixtures were finally fired under inert atmosphere (BOC, Zero Grade Nitrogen <5ppm O₂, flow rate = 5Lmin⁻¹) in a proprietary stainless steel rotary reactor fitted inside a muffle furnace. After reaction the reactor was removed from the muffle furnace and cooled to <50°C by blowing air across the reactor body whilst maintaining the inert atmosphere. The duration of the described cooling process was approximately 20 minutes.

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To evaluate the temperature effect precursor mixtures were fired at temperatures between 700°C and 1120°C. The reaction time was usually between 3 and 5 hours. The produced samples were analysed using X-Ray diffraction (XRD) to assess phase purity. The monoclinic polymorph of LiMnO₂ started to form at 1000°C. The other phases identified by XRD were orthorhombic o-LiMnO₂, tetragonal LiAlO₂, and rock salt Li₂MnO₃. The latter, and the spinel phase LiMn₂O₄, were preferentially formed if the oxygen partial pressure in the reactor exceeded 10⁻⁴atm. Thus, the inert atmosphere should preferably involve a pO₂ of 10⁻⁵ atm or less. This is sufficient to facilitate the formation of manganese in its 3+ oxidation state.

The effect of temperature is summarised in Figure 1.

The highest amounts of the monoclinic polymorph m-LiMnO₂ were obtained at 1060°C, yielding 85% m-LiMnO₂, 5% o-LiMnO₂, 5% of LiAlO₂, and 5% Li₂MnO₃.

It should be noted that this reaction was performed with in-situ grinding by means of a steel rod.

Higher temperature (above 1060°C) appeared to reduce the amount of the monoclinic phase.

More detailed temperature effect results, and X-ray diffraction data are included below under Example II.

Samples of the end product were digested in hydrochloric acid and hydrogen peroxide and subsequently analysed by AA (for Li) and ICP-OES (for Mn and Al). The molar ratio Li:Al:Mn was found to be substantially 1.00:0.05:0.95±5%.

Reaction time was investigated at a constant reaction temperature of 1060°C. The results are shown in Figure 2.

The reaction seems to be completed in approximately three hours. Reaction times of longer than 5 hours usually lead to a decrease of the LiMnO₂-phases and the formation of spinel phase. Reaction time and temperature are likely to be dependent on particle size, and the required reaction may be achieved in a shorter time employing reagents, particularly manganese oxide, having a smaller particle size.

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Example II

In the following example an alumina-lined steel reactor was employed in place of a conventional rotary stainless steel reactor.

Exact stoichiometric amounts (based on LiAl_{0.05}Mn_{0.95}O₂) of lithium carbonate (99.9% pure), manganese (IV) oxide (90% + to <99.0% pure) and aluminium oxide (99.8% pure) were homogenised for 30 minutes using a vibrating table. The resulting mixtures were fired at temperatures between 800°C and 1100°C under inert atmosphere (BOC, zero grade nitrogen) in alumina-lined stainless steel rotary reactors fitted inside a muffle furnace and cooled to less than 50°C by blowing air across the reactor body whilst maintaining the inert atmosphere.

Samples resulting from processing for about 5 hours at 1040°C had average phase compositions of approximately 80% o-LiMnO₂ and 10% m-LiMnO₂.

X-ray diffraction patterns (see Figure 3 A-F) showed a clear shift in the proportion of LiMnO₂ at a temperature of 950°C (compare Figures 3C and D) and higher, and clear identification of the monoclinic form at a temperature of 1050°C (Figure 3F). However, as can be seen from Table 1, the proportion of

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phases was unexpected in comparison with the results of Example I using a stainless steel reactor. Even at 1050°C there was only about 15% m-LiMnO₂.

<u>Table 1:</u> Temperature effect on LiMnO₂ proportions and phases, all at 5 hours reaction time

Temperature °C	Li ₂ MnO ₃	o-LiMnO ₂ %	m-LiMn O ₂	Mn ₃ O ₃
800	35	50	0	15
850	25	65	0	10
900	30	60	0	10
950	5	90	0	5
1000	<5	-95	0	<5
1050	≤5	80	15	ō

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Elemental analysis revealed the presence of high amounts of iron and chromium in samples produced in the conventional steel reactors in comparison with the samples produced in alumina-lined reactors. The Fe content of samples prepared in a steel reactor was generally in the range 1-3%, with Cr in the range 0.05-0.2%. These figures were reduced by about a factor of 10 in the alumina reactor.

It was concluded that a few percent of one or both of Fe and Cr, transferred into the reaction mixture from the steel reactor, aided the formation of the monoclinic LiMnO₂ phase under reaction conditions where it is otherwise difficult to form m-LiMnO₂.

Example III

The process of Example II was repeated in alumina boats in a tube furnace, with the addition of chromium (in the form of Cr₂O₃) to the reaction mixture and subsequent firing at temperatures from 1050 to 1100°C for 3 to 7 hours, under an inert atmosphere. As in Example II, the mixture was first heated to

about 200°C for about an hour before increasing the temperature to the selected level. The results are shown in Table 2.

Table 2: Chromium doping of Aluminium-doped LiMnO2

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Cr ₂ O ₃ Content wt%	Reaction Temp/time °C/h	o-LiMnO₂ %	m-LiMnO ₂	Y-LiAO2 %	LiCrO ₂ %	Li ₂ MnO ₃ %
1	1050/5	5	80	5	0	10
2	1050/5.5	0	95	0	0	5
2.5	1050/7	0	80	5	5	10

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As can be seen from Table 2, the presence of Cr₂O₃ 1 to 2.5 wt%, with a reaction temperature of 1050°C for a period of at least 5 hours, resulted in a marked change in the phase composition of the end product in comparison with the results obtained under similar conditions, but without chromium, in the alumina-lined reactor.

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Whilst these results were obtained using chromium oxide, it is anticipated that other metals (such as Co and Fe) or combinations of metals would similarly facilitate the formation of Al-doped m-LiMnO₂. It is also considered that the chromium, or an alternative metal, may reduce the temperature at which Al-doped m-LiMnO₂ is formed.

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Example IV

The effects of chromium doping were investigated in an alumina-lined reactor using the reagents described for Example II with the addition of chromium oxide (Cr₂O₃), and at 1050°C for 5 hours.

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X-ray diffraction patterns for compositions resulting from the use of different starting proportions of Cr₂O₃ are shown in Figures 4A - D. X-ray diffraction analysis revealed the estimates shown in Table 3 for the content of the compositions.

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<u>Table 3:</u> Chromium doping of aluminium-doped LiMnO₂, in alumina reactor

Cr ₂ O ₃ Content wt%	Reaction Temp/time °C/h	o-LiMnO₂ %	m-LiMnO ₂	Y-LiAIO2 %	LiCrO ₂ %	Li ₂ MnO ₃ %
2	1050/5	40	50	0	.,0	10
3	1050/5	45	35	5	5	5
5	1050/5	15	70	0	10	5
7	1050/5	20	65	0	10	5

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At 3 wt% Cr₂O₃, LiMnO₂ was still predominantly in the orthorhombic form (45%), although there was a 35% proportion of monoclinic form contrasting with only 15% under the same conditions but without chromium doping (see Figure 3). With 5 wt% Cr₂O₃, LiMnO₂ was predominantly in the monoclinic form (70%). The higher proportion of chromium oxide required in this Example compared with Example III could indicate inadequate mixing of the starting materials. Pregrinding of the reaction mixture could significantly improve the effect of chromium.

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Example V

The effects of doping chromium alone in small quantities in LiMnO₂ were systematically investigated in a vertical cylindrical reactor lined with a high

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temperature stable lining. Exact stoichiometric amounts of lithium carbonate (99.9% pure), MnO_2 (99.0% pure) and Cr_2O_3 (99.0% pure) to give products of formula $LiCr_xMn_{1-x}O_2$, where x=0.01, 0.03, 0.05, 0.10, and 0.20, were mixed and ball-milled for 24 hours in a rotating shaker. The resulting mixtures were loaded into the bottom of the reactor and fired at 1000 °C for 5 hours with a flow of zero grade nitrogen percolating through the reaction mixture from an inlet at the bottom of the reactor, with stirring at regular intervals to ensure homogeneity. Elemental analyses of the products using the method described in Example VI confirmed that the end products had the correct Li:Cr:Mn ratios.

The X-ray diffraction patterns of the products are shown in Figure 5. The phase mixtures obtained, estimated from the X-ray diffraction analyses of the products, are given in Table 4. When x is between 0.05 and 0.20 the products are 100% single phase monoclinic $\text{LiCr}_x\text{Mn}_{1-x}\text{O}_2$, as indicated by the data of Davidson et al. in J. Power Sources 54, 205 (1995) and Dahn et al. in J. Electrochem. Soc. 145, 851 (1998). However the results show that predominantly monoclinic $\text{LiCr}_x\text{Mn}_{1-x}\text{O}_2$ can also be obtained when x=0.01 or 0.03, a result not predicted by the aforementioned publications.

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Table 4: Chromium doping of LiMnO2 at 1000 °C

x in LiCr _x Mn _{1-x} O ₂	% orthorhombic phase	% monoclinic phase
0.01	30	70
0.03	10	90
0.05	0	100
0.10	0	100
0.20	0	100

25 The monoclinic layered structure (space group C2/m) of the LiCr_xMn_{1-x}O₂ products was confirmed by Rietveld analysis of the X-ray diffraction data. The unit cell parameters calculated from the X-ray diffraction data by the Rietveld refinement method are shown in Table 5 for the single-phase monoclinic LiCr_xMn_{1-x}O₂ products where x=0.05, 0.10 and 0.20, compared

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with published data for non-doped monoclinic layered LiMnO₂ (Capitaine et al., Solid State Ionics <u>89</u>, 197 (1996)).

Chromium doping results in changes in the crystal unit cell dimensions consistent with substitution of Cr ions into Mn sites of the crystal structure. In LiMnO₂, the extent of the monoclinic crystal distortion resulting from the Jahn Teller distortion of the octahedral coordination environment around the Mn ions is reflected in the magnitude of the ratio between the a and b crystal axes. The systematic decrease in the a/b ratio with increasing amounts of Cr in the crystal is an effect of a minimally distorted octahedral sphere of coordination around the Cr ions. The atomic positions of the cations in the crystal lattice were also refined using the Rietveld method. The results confirmed that Cr and Mn ions share the transition metal sites in the layered crystal structure. No Mn or Cr ions were detected in the interlayer (lithium) sites of the crystal.

<u>Table 5:</u> Unit cell parameters for the single-phase monoclinic LiCr_xMn_{1-x}O₂ products

x in LiCr _x Mn _{1-x} O ₂	a(Å)	b(Å)	c(Å)	β(°)
0.00*	5.439(3)	2.809(2)	5.395(4)	115.9(4)
0.05	5.434(1)	2.809(1)	5.389(2)	115.98(2)
0.10	5.405(2)	2.815(1)	5.375(2)	115.61(2)
0.20	5.382(3)	2.823(1)	5.350(3)	115.05(4)

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Example VI

The effects of using a transitional alumina as the source of Al in the solid state reaction were investigated in a vertical cylindrical reactor with a high

^{*} Capitaine et al. (Solid State Ionics 89, 197 (1996)).

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temperature stable lining. Exact stoichiometric amounts of lithium carbonate (99.9% pure), MnO₂ (99.0% pure) and theta-Al₂O₃ (99.95% pure) to give a product of formula LiAl_{0.05}Mn_{0.95}O₂ were mixed and ball-milled for 30 minutes. The resulting mixture was loaded into the bottom of the reactor and heated to 1000°C for 3.5 hours with a flow of zero grade nitrogen percolating through the reaction mixture from an inlet at the bottom of the reactor, with stirring at regular intervals to ensure homogeneity. After 3.5 hours the product was allowed to cool in the furnace over a period of 15 hours, maintaining the nitrogen flow until the temperature dropped below 50 °C.

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The X-ray diffraction pattern of the product, shown in Figure 6, shows that the product contains approximately 90% m-LiMnO₂, and approximately 10% o-LiMnO₂. No additional impurity phases were detected.

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Samples of the product were digested in hydrochloric acid and hydrogen peroxide and analysed by AA (for Li) and ICP-OES (for other metal ions). The molar ratio Li:Al:Mn was found to be 1.03:0.05:0.94, consistent with the expected stoichiometry within experimental error (±5%) of the analytical technique. Less than 0.1% Cr, 0.1% Ni, and 0.4% Fe were detected in the product, indicating that contamination from the reactor lining was negligible and unlikely to have contributed to the formation of the monoclinic phase.

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The formation of a predominantly monoclinic structure at 1000 °C in this case is attributed mainly to the higher reactivity of the theta-Al₂O₃ by comparison with the aluminium-containing reactants used in the preceding examples. When the reaction was carried out under the same conditions but using an alpha-Al₂O₃ as the source of aluminium, mainly orthorhombic LiMnO₂ was produced. When alpha-Al₂O₃ was used, XRD also detected traces of LiAlO₂ in the product. This was not observed when the theta-Al₂O₃ was used, suggesting that a more complete reaction of the Al₂O₃ with the manganese

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oxide is the critical factor favouring the formation of monoclinic phase under these particular conditions.

The unit cell parameters for the monoclinic phase, obtained by performing a simultaneous Rietveld refinement of the crystal parameters for both the monoclinic and the orthorhombic phase from the X-ray diffraction data, are compared with those given by Capitaine et al. (Solid State Ionics 89, 197 (1996)) for non-doped monoclinic LiMnO₂ in Table 6. The monoclinic LiAl_{0.05}Mn_{0.95}O₂ phase of this example shows a contracted crystal lattice by comparison with the undoped compound. This is consistent with the presence of the Al ion, which has a smaller ionic radius than trivalent Mn, sharing the same crystallographic site as Mn. The contraction is more significant along the a crystal axis than the b crystal axis, consistent with a minimally distorted octahedral sphere of co-ordination around the Al ions since the magnitude of the a/b ratio is related to the average degree of distortion of the octahedron of oxygen ions around the cations in the Mn layer.

<u>Table 6:</u> Unit cell parameters for the monoclinic LiAl_{0.05}Mn_{0.95}O₂ phase of Example VI, compared with reported values for undoped monoclinic LiMnO₂ (Capitaine et al.)

	a(Å)	b(Å)	c(A)	β(°)
LiAl _{0,05} Mn _{0,95} O ₂	5.419(2)	2.802(1)	5.381(2)	115.89(2)
LiMnO ₂ (Capitaine et al.)	5.439(3)	2.809(2)	5.395(4)	115.9(4)

These results demonstrate that the choice of alumina reagent and ball milling of the reactant mixture before heating can influence the temperature at which a predominantly monoclinic form of aluminium doped LiMnO₂ is obtained using the solid state reaction method of the invention.

Electrodes were prepared by mixing 80 wt.% of LiAl_{0.05}Mn_{0.95}O₂, 12 wt.% acetylene black and 8 wt.% poly(vinylidene fluoride) as a slurry in 1-methyl-2-pyrrolidinone (NMP). The slurry was coated onto aluminium foil. After evaporation of the solvent, the coating was pressed on the aluminium foil and

annealed at 150 °C under vacuum. Circular electrodes measuring 14 mm in diameter were then punched from the coated foil. The circular electrodes were weighed individually and the active mass (the total weight of the circular electrode multiplied by the fraction of the electrode weight made up by LiAl_{0.05}Mn_{0.95}O₂) calculated. The electrodes were then dried at 150 °C under vacuum to remove traces of water and transferred to an argon-filled dry glove box (<1 ppm water).

The electrodes were assembled into electrochemical cells within the argon-filled glove box using 2032 button cell hardware. The cell assembly is schematically illustrated in Figure 7. The electrode containing the LiAl_{0.05}Mn_{0.95}O₂ was the cathode 1. The anode 2 was a circular disk of lithium foil having a thickness of 0.38 mm; pressed into a stainless steel lid 3. A polymeric gasket 4 was then positioned over the lip of the lid. A porous glass fibre disk separator 5 wetted with 1M LiPF₆ in (50 wt.% ethylene carbonate + 50 wt.% dimethylene carbonate) electrolyte solution was placed between the anode and cathode. A stainless steel disk 6 and spring 7 were then positioned behind the cathode and the entire assembly hermetically closed within the stainless steel casing 8 by crimp sealing.

Cells prepared by this procedure were cycled more than 200 times between 2.0 and 4.4 V at ambient temperature and elevated temperature (55 °C). A constant charging current of 30 or 75 mA/g was applied until 4.4 V was reached, then the cell was held at 4.4 V until the current dropped below 3 mA/g. Cells were discharged at constant currents of 30 or 75 mA/g. Typical discharge capacities obtained by this procedure at the 30 mA/g discharge rate are given for the 1st, 16th, and 200th cycles in Table 8 in Example VIII. The discharge capacities at ambient temperature increase with cycling. At 55 °C the discharge capacities are particularly high and show good stability.

Example VII

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Exact stoichiometric amounts of lithium carbonate (99.9% pure), MnO₂ (99.0% pure) and theta-Al₂O₃ (99.95% pure) to give a product of formula LiAl_{0.10}Mn_{0.90}O₂ were mixed and ball-milled for 30 minutes. The resulting mixture was loaded into the bottom of the reactor described in Example VI and heated to 1050°C for 10 hours under a flow of zero grade nitrogen. After 10 hours the product was allowed to cool in the furnace over a period of 15 hours, maintaining the nitrogen flow until the temperature dropped below 50°C.

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The X-ray diffraction pattern of the product, shown in Figure 8, shows that the product contains approximately 95% m-LiMnO₂, with the remainder being o-LiMnO₂ phase. No additional impurity phases were detected.

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The unit cell parameters of the monoclinic phase, calculated from the XRD pattern, are a = 5.416(2) Å, b = 2.804(1) Å, c = 5.382(2) Å, and $\beta = 115.91^{\circ}$. Like the monoclinic LiAl_{0.05}Mn_{0.95}O₂ phase of Example VI, the LiAl_{0.10}Mn_{0.90}O₂ material of this example shows a contracted crystal lattice by comparison with the undoped compound.

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Electrodes and electrochemical cells containing the LiAl_{0.10}Mn_{0.90}O₂ material were prepared as described in Example VI, and the cells were charged and discharged following the same procedures. Typical discharge capacities obtained at the 30 mA/g discharge rate are given for the 1st, 16th, and 200th cycles at ambient and 55 °C in Table 8 in Example VIII. The material shows slightly lower initial capacities than the LiAl_{0.05}Mn_{0.95}O₂ compound of Example VI, but the capacity increases with cycling to values which are similar to those of the LiAl_{0.05}Mn_{0.95}O₂ material.

Example VIII

Compositions LiAl_xCr_yMn_{1-x-y}O₂ were prepared by reacting lithium carbonate (99.9% pure), MnO₂ (99.0% pure), theta-Al₂O₃ (99.95% pure) and Cr₂O₃ (99.0% pure). Exact stoichiometric amounts of the reactants to give the compositions LiAl_{0.05}Cr_{0.01}Mn_{0.94}O₂, LiAl_{0.05}Cr_{0.03}Mn_{0.92}O₂, LiAl_{0.05}Cr_{0.05}Mn_{0.93}O₂ were mixed and ball-milled for 24 hours in a rotating shaker, then loaded into the vertical cylindrical reactor of Example VI and heated at 1050 °C for 5 hours under a flow of zero grade nitrogen.

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The X-ray diffraction patterns of the products are shown in Figure 9(A-D). In the case of the compositions LiAl_{0.05}Cr_{0.01}Mn_{0.94}O₂ and LiAl_{0.05}Cr_{0.03}Mn_{0.92}O₂, the products were 80-90 % layered monoclinic phase, with the remaining phase being o-LiMnO₂. No other impurity phases were observed. The compositions LiAl_{0.05}Cr_{0.05}Mn_{0.90}O₂ and LiAl_{0.02}Cr_{0.05}Mn_{0.93}O₂ were phase-pure layered monoclinic materials. Cell parameters for these phases are given in Table 7.

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<u>Table 7:</u> Unit cell parameters for the monoclinic LiAl_xCr_yMn_{1-x-y}O₂ products

x and y in	. a	b	С	β
LiAl _x Cr _y Mn _{1-x-y} O ₂	(Å)	(Å)	(Å)	(°)
x = 0.05, y = 0.01	5.424	2.805	5.394	115.87
x = 0.05, y = 0.03	5.426	2.809	5.390	115.90
x = 0.05, y = 0.05	5.408	2.808	5.375	115.77
x = 0.03, y = 0.05	5.415	2.810	5.379	115.78

Electrodes and electrochemical cells containing the LiAl_{0.05}Cr_{0.05}Mn_{0.90}O₂, and LiAl_{0.02}Cr_{0.05}Mn_{0.93}O₂ materials were prepared as described in Example VI, and the cells were charged and discharged following the same procedures. Typical discharge capacities obtained at the 30 mA/g discharge rate are given for the 1st, 16th, and 200th cycles at ambient temperature and 55 °C in Table 8. The cells cycled at 55 °C showed similar excellent capacity retention over 200 cycles as the LiAl_{0.05}Mn_{0.95}O₂ and LiAl_{0.10}Mn_{0.90}O₂ materials of Examples VI and VII. However the LiAl_{0.05}Cr_{0.05}Mn_{0.90}O₂, and LiAl_{0.02}Cr_{0.05}Mn_{0.93}O₂ materials showed increased overall capacity at both 55 °C and ambient temperature when compared with the LiAl_{0.05}Mn_{0.95}O₂ and LiAl_{0.10}Mn_{0.90}O₂. materials.

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The combination of chromium plus aluminium together in monoclinic layered LiMnO₂ therefore provides a material with improved cycling characteristics over monoclinic layered LiMnO₂ doped with aluminium alone. The addition of chromium to the aluminium-doped LiMnO₂ also facilitated the obtaining of a single-phase monoclinic layered material.

<u>Table 8:</u> Discharge capacities for the monoclinic LiAl_xMn_{1-x}O₂ materials of Examples VI, VII & IX, and the LiAl_xCr_yMn_{1-x-y}O₂ materials of Example VIII.

		Discharge capacity (mAh/g) at 30 mA/g current rate 4.4-2 V						
Material	Example	Ambient temperature			55 °C			
		Cycle	Cycle Cycle Cycle		Cycle	Cycle	Cycle	
Tel.		1	16	196	1	16	196	
LiAl _{0.05} Mn _{0.95} O ₂	VI	82	82	139	156	164	168	
LiAl _{0.05} Mn _{0.95} O ₂	X	92	92	140	161	177	152	
LiAl _{0.10} Mn _{0.90} O ₂	VII	73	73	143	137	162	168	
LiAl _{0.05} Cr _{0.05} Mn _{0.90} O ₂	VIII	93	80	127	158	182	184	
LiAl _{0.02} Cr _{0.05} Mn _{0.93} O ₂	VIII	122	138	122	170	184	170	

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Example IX

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Exact stoichiometric amounts of lithium carbonate (99.9% pure), manganese carbonate (99.5% pure) and Cr_2O_3 (99.0% pure) to give a product of formula $Li_{0.5}Cr_{0.05}Mn_{0.95}O_2$ were mixed and ground in an acetone slurry using a mortar and pestle. The mixture was loaded into a vessel formed from an inert metal alloy and fired in a furnace at 700 °C for 20 hours in air. The X-ray diffraction pattern of the product, in Figure 10A, shows that it was a single phase cubic spinel with a unit cell parameter of a = 8.235 Å. The X-ray diffraction data are consistent with doping of the chromium on the manganese site of spinel $LiMn_2O_4$.

This compound was ground in acetone with an equivalent of lithium carbonate to give a product of formula LiCr_{0.05}Mn_{0.95}O₂, and heated under a flow of zero grade nitrogen at 1000 °C for 18 hours. The X-ray diffraction pattern of this product, in Figure 10B, shows formation of phase-pure monoclinic layered LiMnO₂, with unit cell parameters a = 5.431 Å, b = 2.807Å, c = 5.382 Å, and $\beta = 115.94$ °. These cell parameters are very close to those obtained for the LiCr_{0.05}Mn_{0.95}O₂ compound of Example V.

This example shows that a two stage solid state process, in which a doped spinel phase is formed by reaction in air and then reacted with additional lithium under nitrogen, can be used to synthesise doped LiMnO₂ materials. This reaction route provides an alternative to the single-step reaction, and may be useful when employing starting oxides or salts (as the source of lithium, manganese or dopant) which are difficult to react together to give a single-phase product under an inert atmosphere. Such oxides or salts may react more easily in air to give a doped oxidised lithium manganese oxide product, which can then be converted into the doped LiMnO₂ in inert atmosphere in a second stage.

It will be appreciated by those skilled in the art that doped oxidised lithium manganese oxide compounds can be formed by solid state reactions at temperatures ranging from 200 °C up to 900 °C, and that a range of temperature conditions could therefore be used in the first stage of this process. The second, conversion step in inert atmosphere should preferably be performed at a higher temperature than the first step.

Example X

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The two stage reaction process was used to prepare Al-doped LiMnO₂. Exact stoichiometric amounts of lithium carbonate (99.9% pure), MnO₂ (99.0% pure) and theta-Al₂O₃ (99.95% pure) to give a product of formula LiAl_{0.05}Mn_{0.95}O₂ were mixed and ball-milled for 30 minutes. The resulting mixture was heated at 700°C in air for 8 hours, cooled, then fired at 1000 °C for 5 hours under a flow of zero grade nitrogen. The product was cooled over a period of one hour, maintaining the nitrogen flow until the temperature dropped below 50 °C.

X-ray diffraction of the product after the first stage of the reaction in air (Figure 11A) showed a mixture of cubic spinel phase and Li₂MnO₃. The presence of Li₂MnO₃ in this case is due to the amount of lithium, one mole of lithium per mole of manganese + aluminium, being too high for a single-phase spinel. The X-ray diffraction pattern of the product after the second stage under nitrogen (Figure 11B) shows complete conversion of the spinel and Li₂MnO₃ phases to LiMnO₂ product. The product is approximately 90% monoclinic layered LiMnO₂, with around 10% o-LiMnO₂, practically identical to the LiAl_{0.05}Mn_{0.95}O₂ product of Example VI.

Electrodes and electrochemical cells containing the LiAl_{0.05}Mn_{0.95}O₂ material of this example were prepared as described in Example V, and the cells were charged and discharged following the same procedures. Typical discharge capacities obtained at the 30 mA/g discharge rate are given for the 1st, 16th,

and 200th cycles at ambient and 55 °C in Table 8 in Example VIII. The electrochemical characteristics are almost identical to those of the LiAl_{0.05}Mn_{0.95}O₂ product of Example VI.

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Example XI

The two-stage synthesis procedure described in Example IX was used to prepare a compound of stoichiometry LiGa_{0.10}Mn_{0.90}O₂. Exact stoichiometric amounts of lithium carbonate (99.9% pure), manganese carbonate (99.5% pure) and gallium oxide Ga₂O₃ (99.9% pure) to give a product of formula Li_{0.5}Ga_{0.10}Mn_{0.90}O₂ were mixed and ground in an acetone slurry using a mortar and pestle. The mixture was loaded into a vessel formed from an inert metal alloy and fired in a furnace at 700 °C, for 20 hours in air. The X-ray diffraction pattern of the product, in Figure 12A, shows that it was a single phase cubic spinel with a unit cell parameter of a = 8.227 Å.

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This compound was ground in acetone with an equivalent of lithium carbonate to give a product of formula LiGa_{0.10}Mn_{0.90}O₂, and heated under a flow of zero grade nitrogen at 1000 °C for 15 hours. The X-ray diffraction pattern of this product, in Figure 12B, shows 80% formation of monoclinic layered LiMnO₂, with unit cell parameters a = 5.430 Å, b = 2.810 Å, c = 5.387 Å, and $\beta = 115.78$ °. The remaining 20% of the product is o-LiMnO₂.

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This result demonstrates that gallium may be used as an alternative, or in addition, to aluminium or chromium as a means of stabilising the layered monoclinic modification of LiMnO₂.

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Example XII

The two-stage synthesis procedure described in Example IX was used to prepare compounds of stoichiometry LiCr_{0.05}Ga_{0.05}Mn_{0.90}O₂ and LiCr_{0.05}B_{0.05}Mn_{0.90}O₂. Exact stoichiometric amounts of lithium carbonate (99.9% pure), manganese carbonate (99.5% pure), Cr₂O₃ (99.9% pure) and gallium oxide Ga₂O₃ (99.9% pure) or boric acid H₃BO₃ (99.9% purity) to give products of formulae Li_{0.5}Cr_{0.05}Ga_{0.05}Mn_{0.90}O₂ and Li_{0.5}Cr_{0.05}B_{0.05}Mn_{0.90}O₂ were mixed and ground in an acetone slurry using a mortar and pestle. The mixtures were loaded into a vessel formed from an inert metal alloy and fired in a furnace at 700 °C for 20 hours in air. The X-ray diffraction patterns of the products in Figure 13A&C show that both were single phase cubic spinels with a unit cell parameter of a = 8.227 Å for the Li_{0.5}Cr_{0.05}Ga_{0.05}Mn_{0.90}O₂ material and a = 8.234 Å for the Li_{0.5}Cr_{0.05}B_{0.05}Mn_{0.90}O₂ material.

These compounds were each ground in acetone with an equivalent of lithium carbonate to give products of formulae LiCr_{0.05}Ga_{0.05}Mn_{0.90}O₂ and LiCr_{0.05}B_{0.05}Mn_{0.90}O₂, and heated under a flow of zero grade nitrogen at 1000° C for 15 hours. The X-ray diffraction patterns of the products in Figure 13B&D show the formation of phase-pure monoclinic layered LiMnO₂ in both cases, with unit cell parameters a = 5.410 Å, b = 2.815 Å, c = 5.378 Å, and $\beta = 115.79$ ° for the LiCr_{0.05}Ga_{0.05}Mn_{0.90}O₂ material, and a = 5.429 Å, b = 2.811 Å, c = 5.380 Å, and $\beta = 115.86$ ° for the LiCr_{0.05}B_{0.05}Mn_{0.90}O₂ material.

The above examples show that monoclinic layered LiMnO₂ may be formed during a high temperature solid state reaction involving metal doping, under specific conditions of time and temperature, and with appropriate starting reagents. Zero grade nitrogen is sufficient to control the manganese oxidation state in the formation of the monoclinic LiMnO₂ material, as only very little Li₂MnO₃ and no spinel phases were usually formed in the reactions under zero grade nitrogen.

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The temperature window for producing layered monoclinic LiMnO $_2$ is in the range 800 to 1080°C.

The formation of aluminium doped layered monoclinic LiMnO₂ from lithium carbonate, manganese oxide and aluminium oxide (or hydroxide) precursors by high temperature solid state reaction is facilitated by the inclusion of an additional metal, such as chromium.

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It will be appreciated by those skilled in the art that the selection of reagents, in particular their average particle sizes and reactivity, will influence the optimum conditions of time and/or temperature to achieve the preferred result. As shown in the examples, under appropriately selected conditions, the high temperature solid state reaction method of the invention can produce stable layered monoclinic LiMnO₂ doped with aluminium, chromium, gallium, or combinations of aluminium, chromium, gallium and boron. Combinations of aluminium and chromium result in improved electrochemical properties over materials doped with aluminium alone. It is anticipated that other combinations may also result in improvements.

Where in the foregoing description reference has been made to specific components or integers of the invention having known equivalents then such equivalents are herein incorporated as if individually set forth.

Although this invention has been described by way of example and with reference to possible embodiments thereof it is to be understood that modifications or improvements may be made thereto without departing from the scope or spirit of the invention.

CLAIMS

- 1. A metal-doped lithium manganese oxide composition having the formula $\operatorname{LiM^1_{m_1}M^2_{m_2}...M^x_{m_x}Mn_zO_2}$ wherein $\operatorname{M^1, M^2_{...M}}^x$ are each different metal ions, $\operatorname{ml+m2+...mx+z}$ is substantially equal to 1, $\operatorname{ml+m2+...mx}$ is greater than zero but less than 0.75 and z and at least ml are greater than zero, providing that where $\operatorname{m2...mx}$ are zero $\operatorname{M^1}$ is Cr and $\operatorname{0<ml}<0.05$ or $\operatorname{M^1}$ is Ga .
- 2. A composition according to claim 1 in predominantly monoclinic form.

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- 3. A composition according to either claim 1 or claim 2 wherein M¹, M²...M^x are each selected to have an ionic radius in their relevant oxidation state, similar to or less than that of Mn³⁺.
- 4. A composition according to claim 3 wherein M¹, M²...M^x are each selected from the group consisting of Al, Ga, Cr, Co, Fe, B, V, Ti and Ru.
 - 5. A composition according to claim 4 wherein M¹ is Al.
- 20 6. A composition according to claim 5 wherein M² is selected from Ga, Cr, Co, Fe, B, V, Ti and Ru.
 - 7. A composition according to claim 6 wherein M² is Cr.
- 25 8. A composition according to claim 1 wherein M¹, M²...M^x each has the property of existing in a minimally distorted octahedral sphere of coordination.
- 9. A composition according to claim 1 wherein M¹, M²...M^x each has a formal oxidation state of 3+.

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- 10. A composition according to claim 1 wherein $m_1+m_2+...m_x$ is greater than zero but less than 0.25.
- 11. A composition according to claim 1 of the formula LiAl_{m1}Cr_{m2}Mn_zO₂,

 predominantly in monoclinic form, wherein 0<m1≤0.05, 0<m2≤0.05 and

 0.90≤ z<1.0.
 - 12. A composition according to claim 11 wherein m1=0.05, m2=0.05 and z=0.90.

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- 13. A composition according to claim 11 wherein m1=0.02, m2=0.05 and z=0.93.
 - 14. A composition according to claim 1 of the formula LiCr_{m1}Mn_zO₂;

 predominantly in monoclinic form, wherein 0<m1<0.05 and 0.95<z<1.0.
- 15. A composition according to claim 14 wherein m1=0.03 and z=0.97.
 - 16. A composition according to claim 14 wherein m1=0.01 and z=0.99.
- 20 17. A composition according to claim 1 of the formula $LiGa_{m1}Mn_zO_2$, predominantly in monoclinic form, wherein $0 < m1 \le 0.10$ and $0.9 \le z < 1.0$.
- 18. A composition according to claim 1 of the formula LiM¹_{m1}Cr_{m2}Mn_zO₂
 wherein M¹ is selected from Ga or B, m1 and m2 are each greater than zero
 but less than 0.1 and 0.9<z<1.0.
 - 19. A method of synthesising a metal-doped lithium manganese oxide composition having the formula LiM¹_{m1} M²_{m2}...M^x_{mx} Mn_z O₂ wherein M¹, M²...M^x are each different metal ions, m1+m2+...mx + z is substantially equal to 1, m1+m2+...mx is greater than zero but less than 0.75, and z and at least

m1 are each greater than zero, providing that where m2...mx are zero and M¹ is Cr then 0<m1<0.05, in stable monoclinic form by solid state reaction.

20. A method according to claim 19, wherein the solid state reaction includes:

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- mixing reagents including a lithium compound, a manganese compound and at least one additional metal compound, each compound selected from an oxide, a hydroxide and/or a carbonate, and
- heating the mixture in a reactor under inert atmosphere, to a temperature exceeding 800°C for at least 2 hours.
 - 21. A method according to claim 20 wherein the lithium compound is lithium carbonate.

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- 22. A method according to either claim 20 or claim 21 wherein the manganese compound is manganese oxide.
- 23. A method according to claim 20 wherein the or each metal of the at least one additional metal compound is selected to have a ionic radius in its relevant oxidation state, similar to or less than that of Mn³⁺.
 - 24. A method according to claim 23 wherein the or each metal of the at least one additional metal compound is selected from the group consisting of Al, Ga, Cr, Co, Fe, B, V, Ti and Ru.
 - 25. A method according to claim 24 wherein the at least one additional metal compound includes aluminium oxide and chromium oxide.
- 30 26. A method according to claim 25 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiAl_xCr_yMn_z0₂

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wherein x+y+z=1 and x and y are each ≥ 0.01 but ≤ 0.05 and z is ≥ 0.90 and ≤ 0.98 .

- 27. A method according to claim 24 wherein the at least one additional metal compound includes aluminium oxide.
 - 28. A method according to claim 27 wherein the aluminium oxide is a transitional alumina.
- 10 29. A method according to claim 27 or claim 28 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiAl_xMn_yO₂ wherein x+y=1, 0.01≤x≤0.10 and 0.90≤y≤0.99.
- 30. A method according to claim 24 wherein the at least one additional metal compound includes chromium oxide.
 - 31. A method according to claim 30 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiCr_xMn_yO₂, wherein x+y=1, 0.01≤x<0.05 and 0.95<y≤0.99.
 - 32. A method according to claim 20 wherein the or each metal of the at least one additional metal compound has a formal oxidation state of 3⁺.
- 33. A method according to any one of claims 20 to 32 further including the step 25 of ball-milling the mixed reagents.
 - 34. A method according to any one of claims 20 to 33 wherein the mixture is heated to a temperature in the range substantially 900°C to 1080°C.
- 35. A method according to claim 34 wherein the mixture is heated to a temperature of substantially 1000°C to 1080°C for a period of at least 5 hours.

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36. A method according to any one of claims 20 to 35 wherein the inert atmosphere has a sufficiently low pO₂ to facilitate the formation of manganese in its 3+ oxidation state.

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37. A method according to claim 36 wherein the pO₂ is substantially 10⁻⁵ atm or less.

38. A method according to claim 20 wherein the mixture is heated in a reactor having a high temperature stable lining.

39. A method of synthesising a metal-doped lithium manganese oxide composition of the formula LiM¹_{m1}M²_{m2...}M^x_{mx}Mn_zO₂ wherein M¹, M²...M^x are each different metal ions, m1+m2+...mx+z is substantially equal to 1, m1+m2+...mx is greater than zero but less than 0.75 and z and at least m1 are each greater than zero, in stable predominantly monoclinic form, the method including the steps of:

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- mixing reagents including a lithium compound, a manganese compound and at least one additional metal compound, each compound selected from an oxide, a hydroxide and/or a carbonate,
- heating the mixture in air to a temperature ≥ 200°C for at least 5 hours to produce oxygen-rich lithium manganese oxides,

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- further heating the mixture in a reactor under inert atmosphere to a temperature exceeding 800°C for at least 2 hours.
- 40. A method according to claim 39 wherein the lithium compound is lithium carbonate.

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- 41. A method according to either claim, 39 or claim 40 wherein the manganese compound is manganese carbonate.
- 42. A method according to claim 41 wherein the or each metal of the at least one additional metal compound is selected to have an ionic radius in its relevant oxidation state similar to or less than that of Mn³⁺.
 - 43. A method according to claim 42 wherein the or each additional metal compound is selected from Al, Ga, Cr, Co, Fe, B, V, Ti and Ru.

44. A method according to claim 43 wherein the or each additional metal compound includes at least one of an oxide of Cr, Al, Ga and B.

- 45. A method according to claim 39 wherein the or each additional metal compound has a formal oxidation state of 3⁺.
 - 46. A method according to any one of claims 39 to 45 wherein the heating in air is at a temperature in the range 600 to 850°C.
- A method according to claim 46 wherein the heating in air is at substantially 700°C.
 - 48. A method according to any one of claims 39 to 47 wherein the heating under inert atmosphere is at a temperature of substantially 1000-1080°C.
 - 49. A method according to claim 48 wherein the inert atmosphere has a sufficiently low pO₂ to facilitate the formation of manganese in its 3+ oxidation state.
- 30 50. A method according to claim 48 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiCr_{0.05}Mn_{0.95}O₂.

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- 51. A method according to claim 48 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiAl_{0.05}Mn_{0.95}O₂.
- 5 52. A method according to claim 48 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiM¹_{0.05}Cr_{0.05}Mn_{0.90}O₂ wherein M¹ is selected from Al, Ga or B.
- 53. A method according to claim 48 wherein the reagents are mixed in substantially stoichiometric amounts for the synthesis of LiGa_{0.1}Mn_{0.9}O₂.
 - An electrode material for a battery, said material including a composition according to any one of claims 1 to 18.
- 15 55. A non-aqueous electrolyte lithium secondary battery including a positive electrode, a negative electrode capable of absorbing and desorbing lithium in a reversible manner and a lithium ion conducting non-aqueous electrolyte, wherein the positive electrode includes a composition according to any one of claims 1 to 18.
 - 56. Use of a composition according to any one of claims 1 to 18 in the manufacture of a non-aqueous electrolyte lithium secondary battery.
- 57. A metal-doped lithium manganese oxide composition substantially as herein described and with reference to the accompanying Examples.
 - 58. A method of synthesising a metal-doped lithium manganese oxide composition substantially as herein described and with reference to the accompanying Examples.

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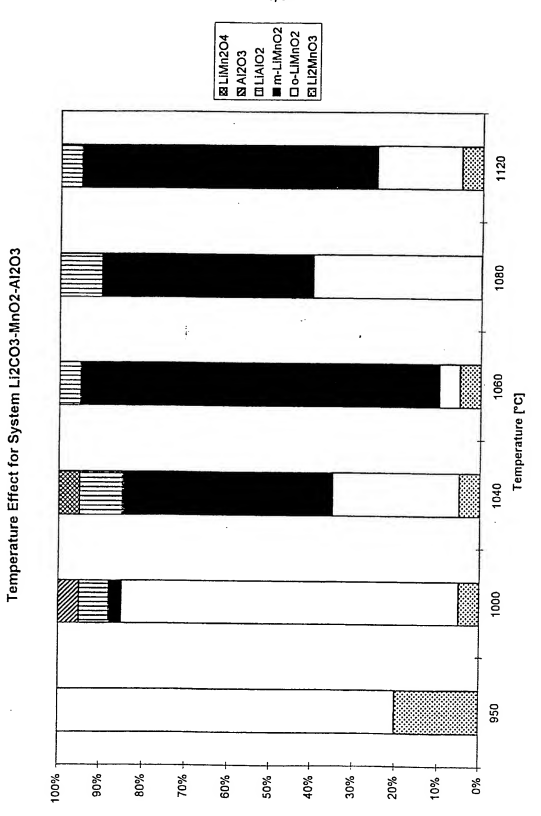
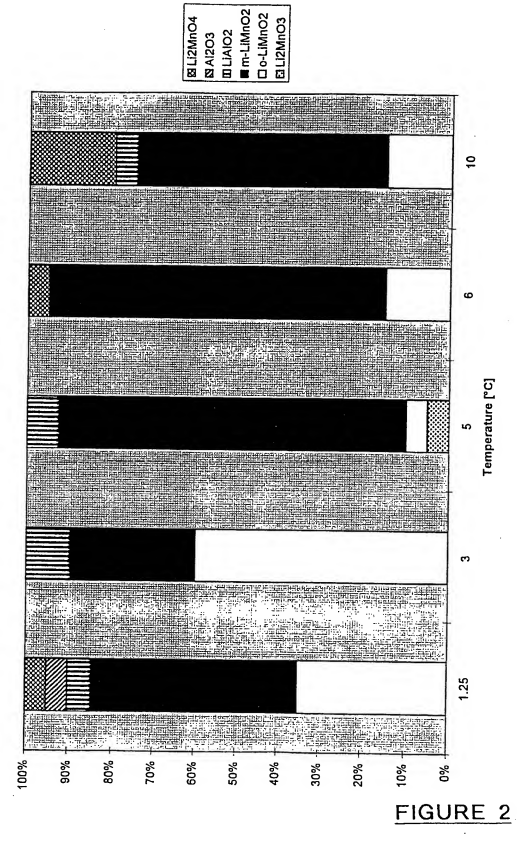
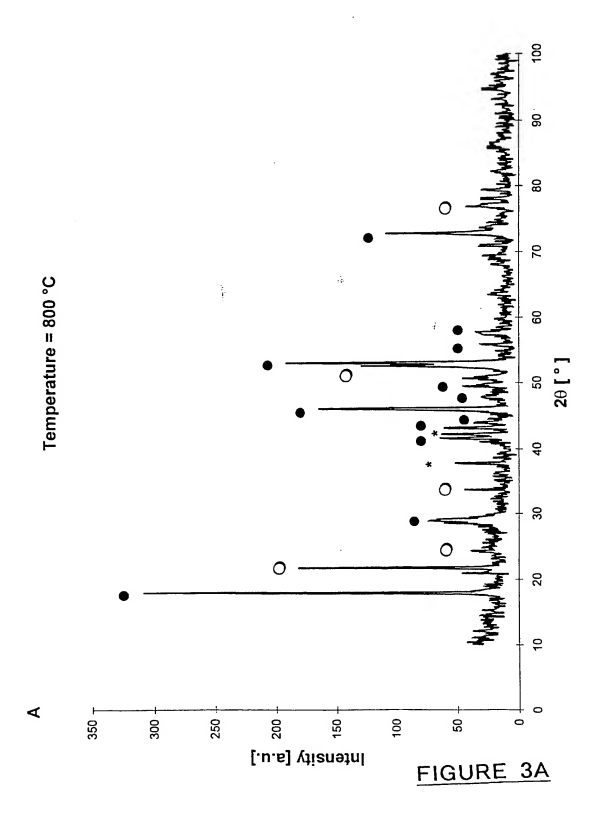
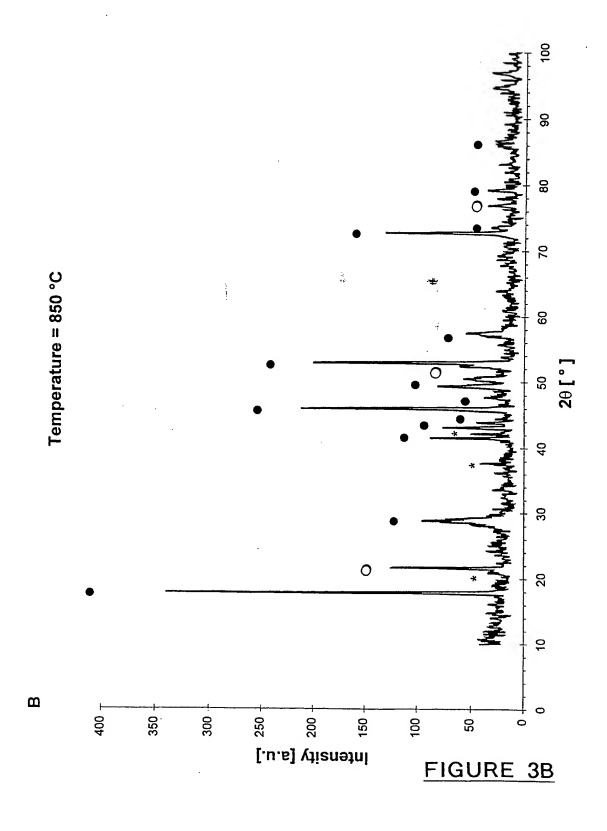


FIGURE 1

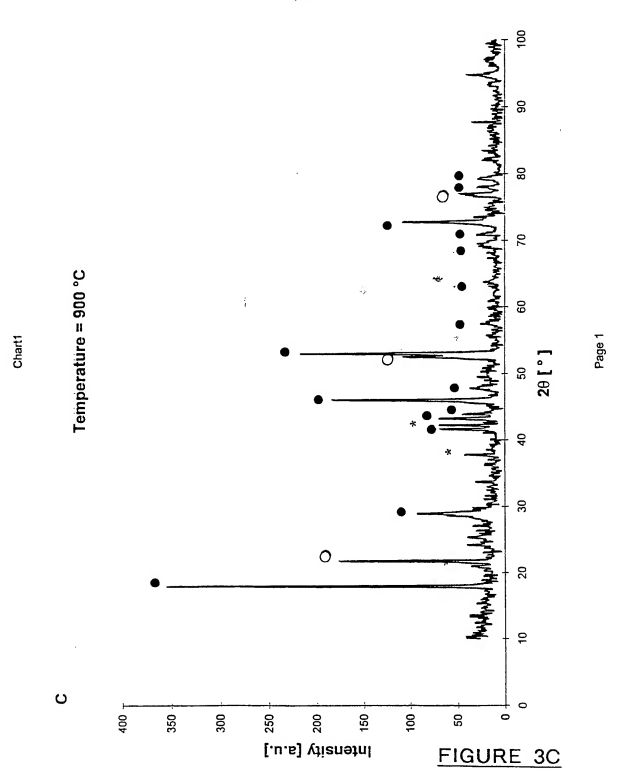


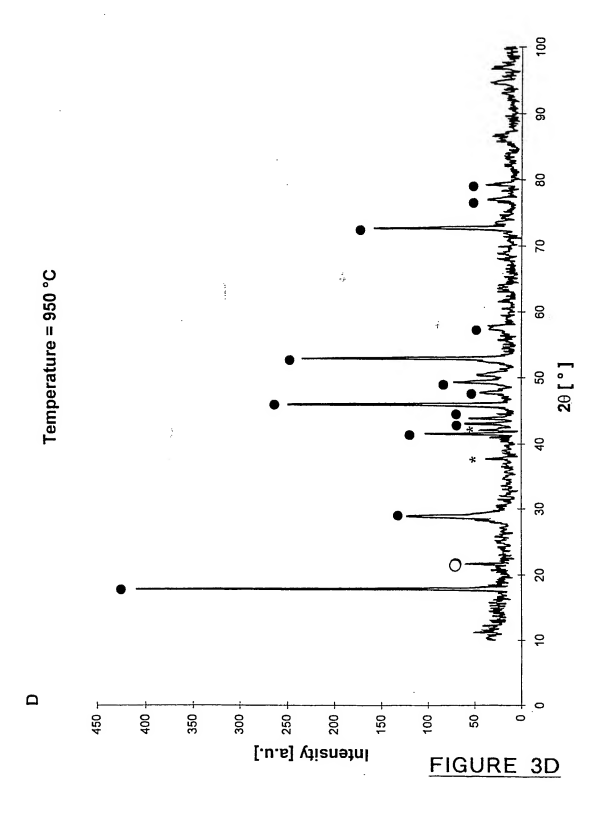


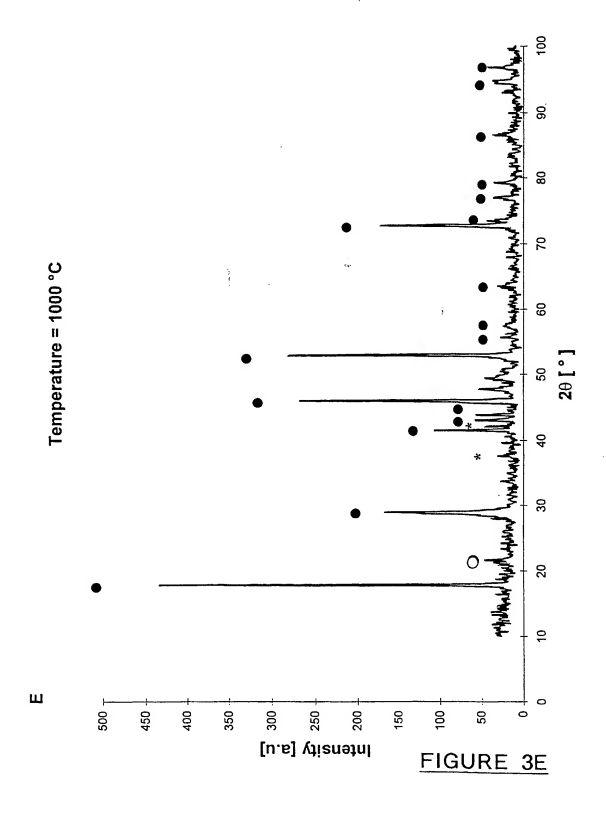


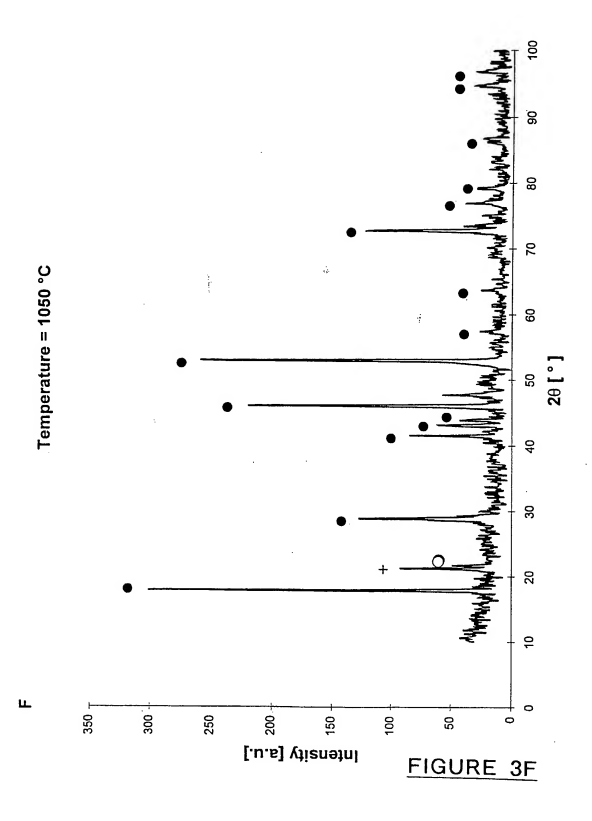


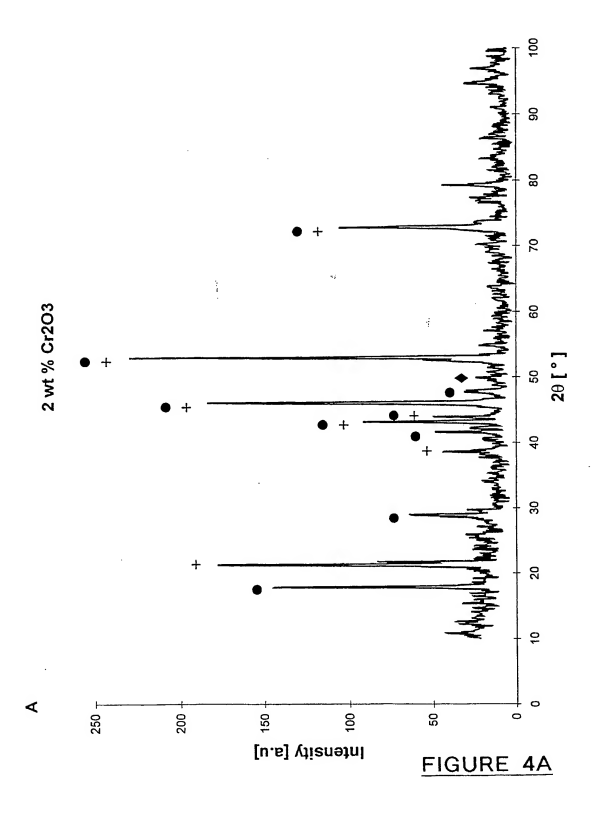


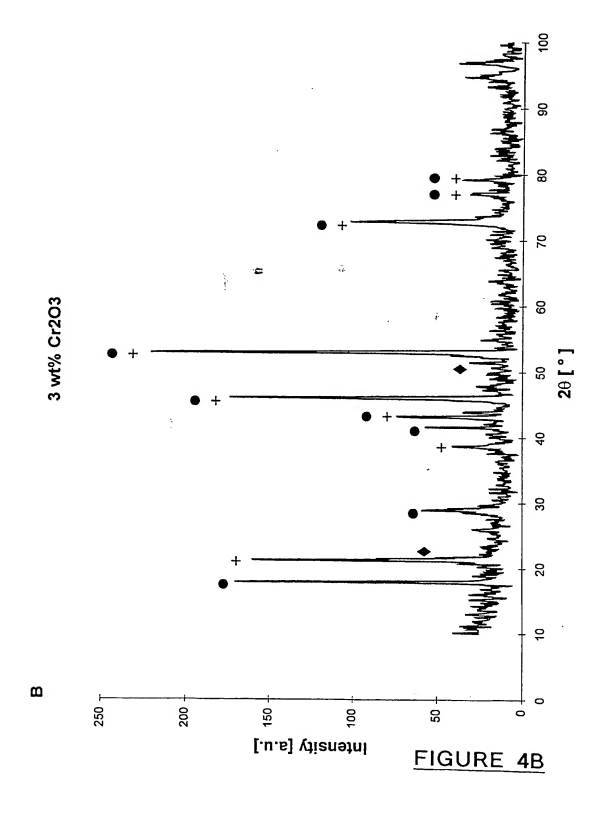


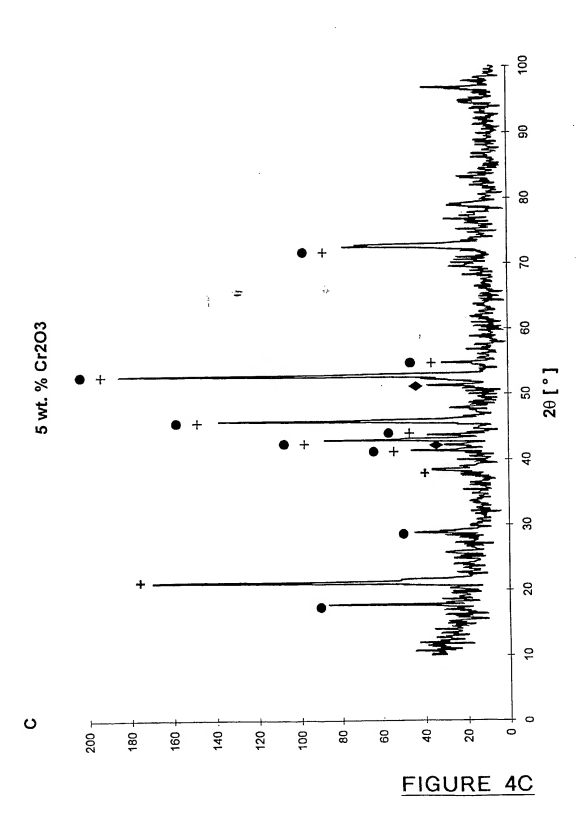


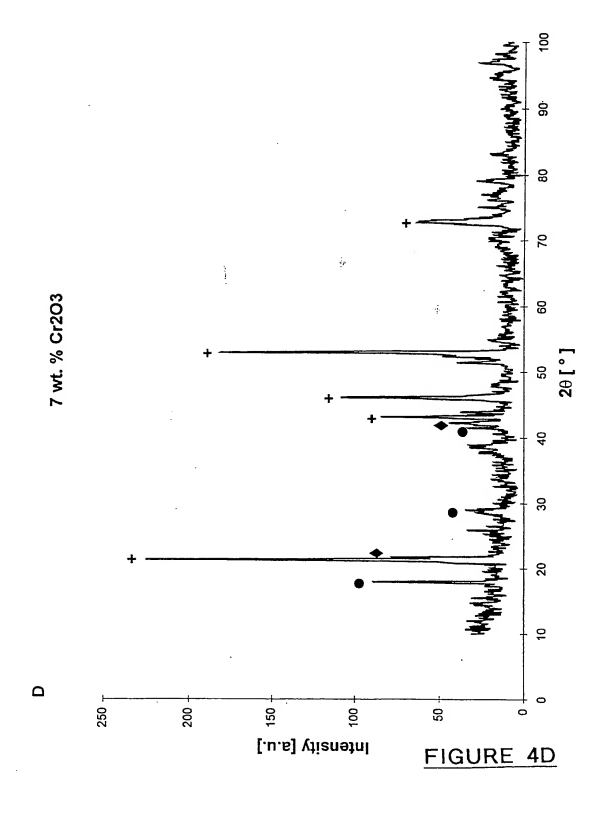












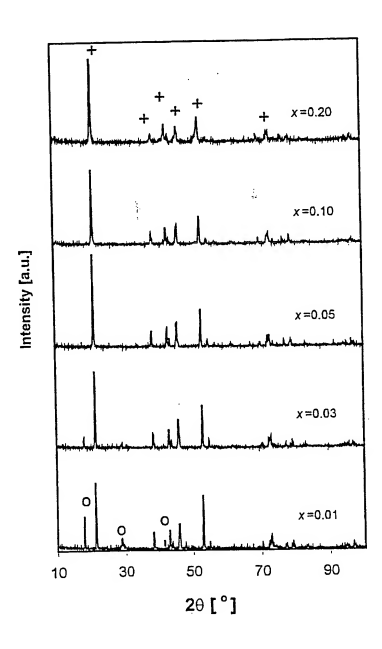


FIGURE 5

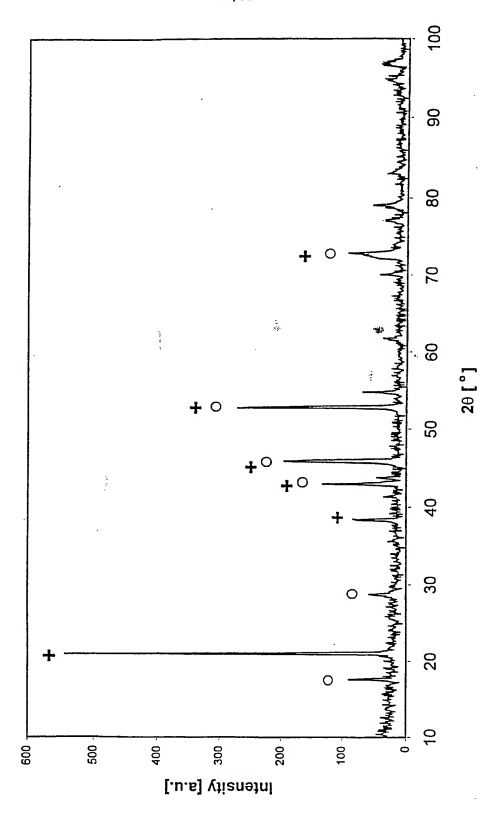
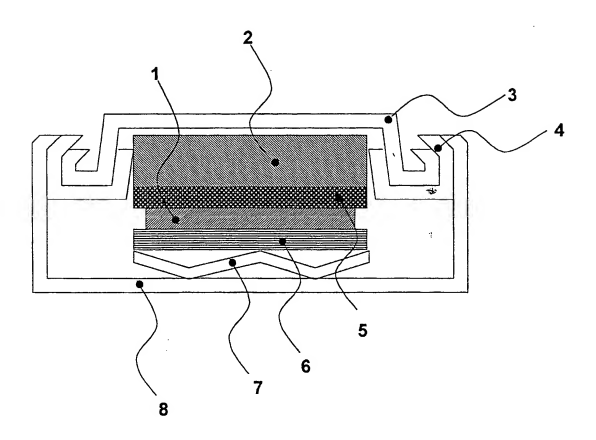


FIGURE 6



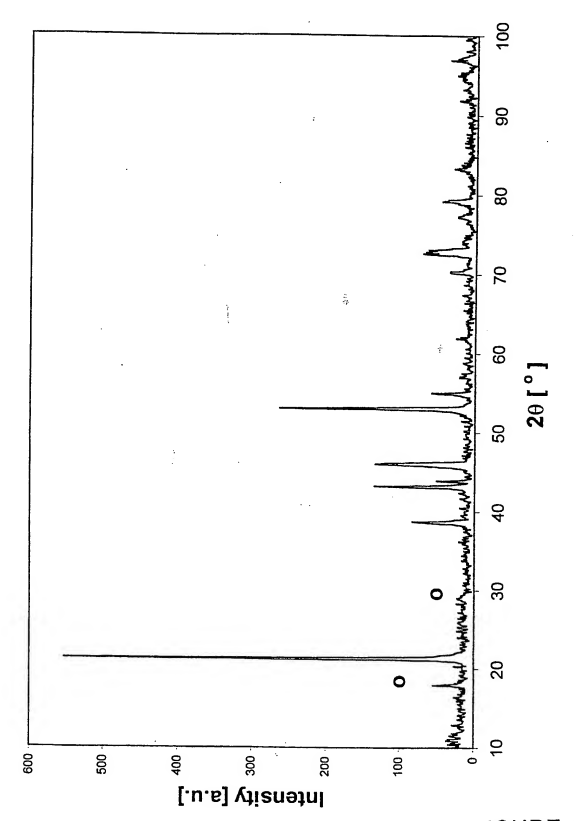


FIGURE 8

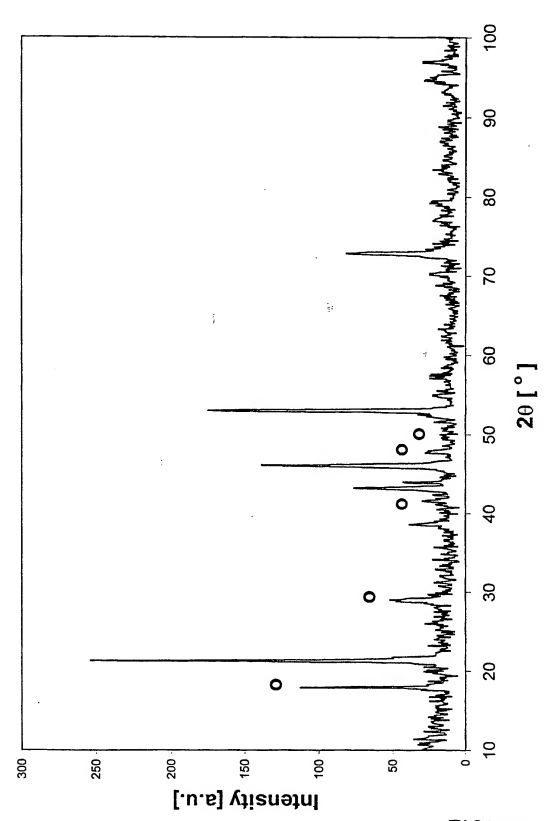
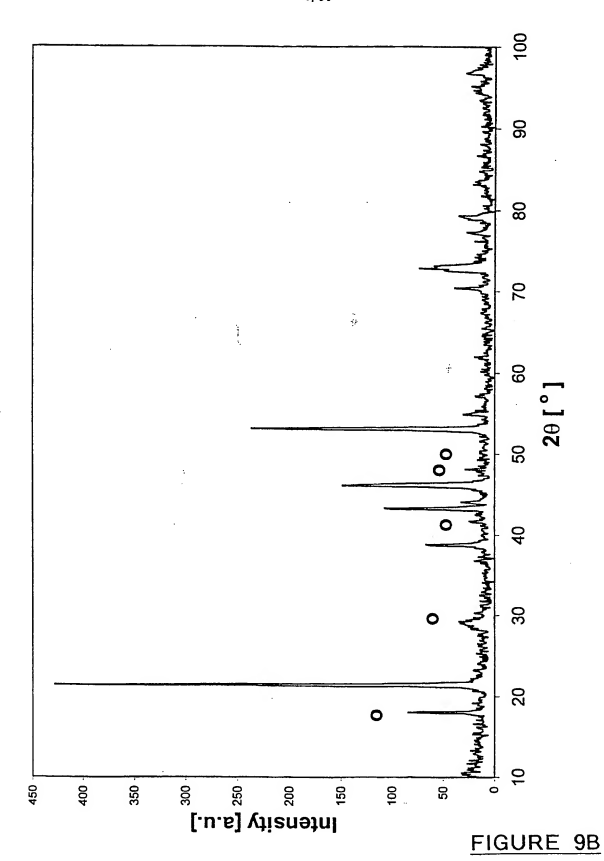


FIGURE 9A



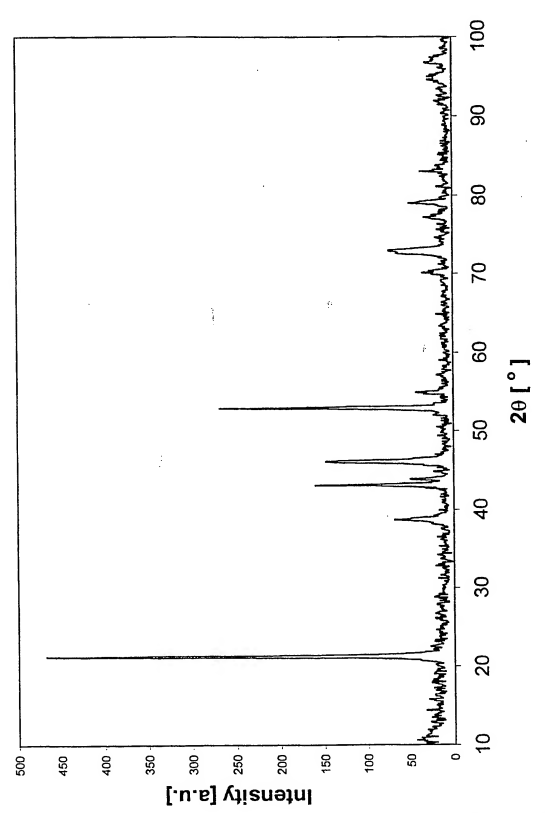


FIGURE 9C

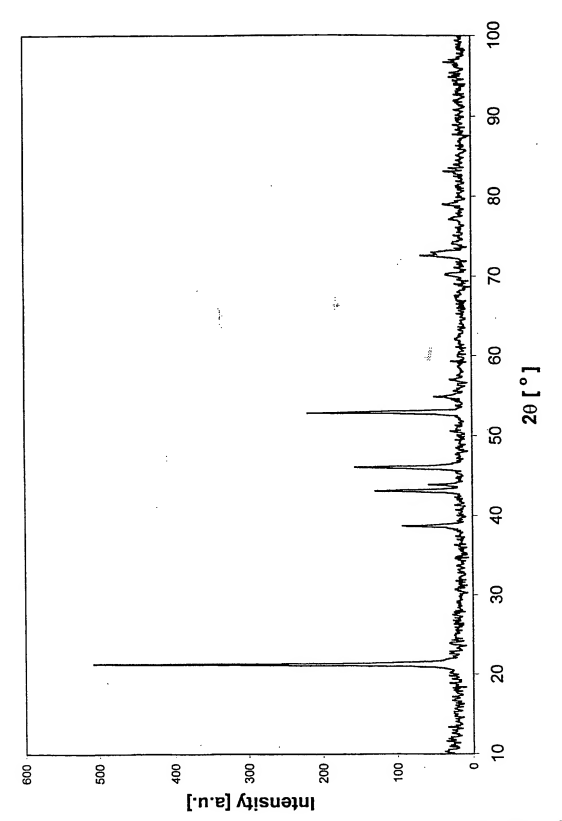


FIGURE 9D

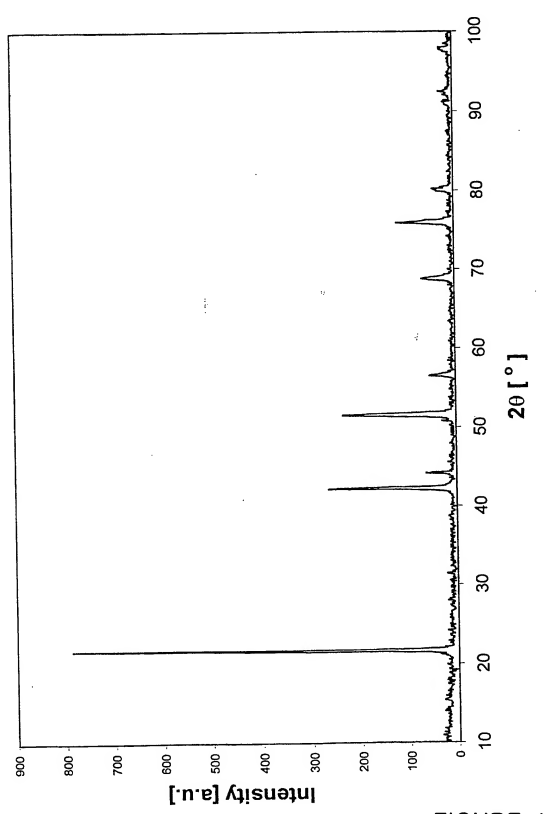


FIGURE 10A

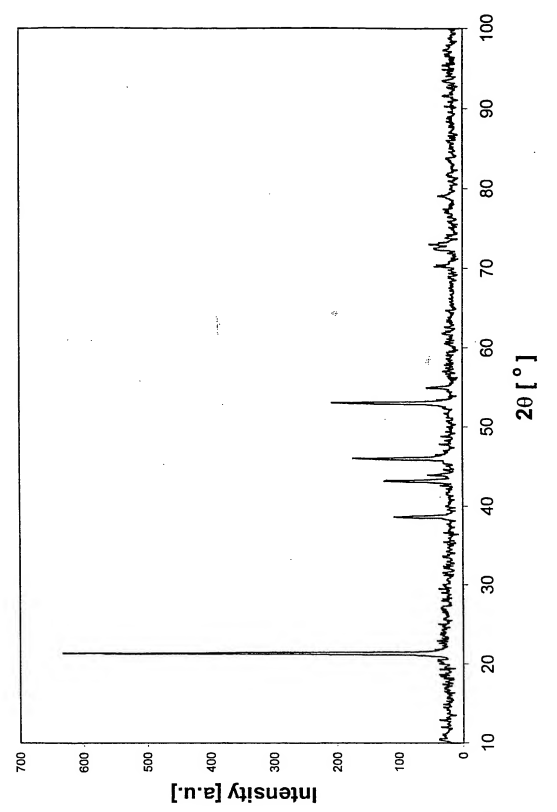
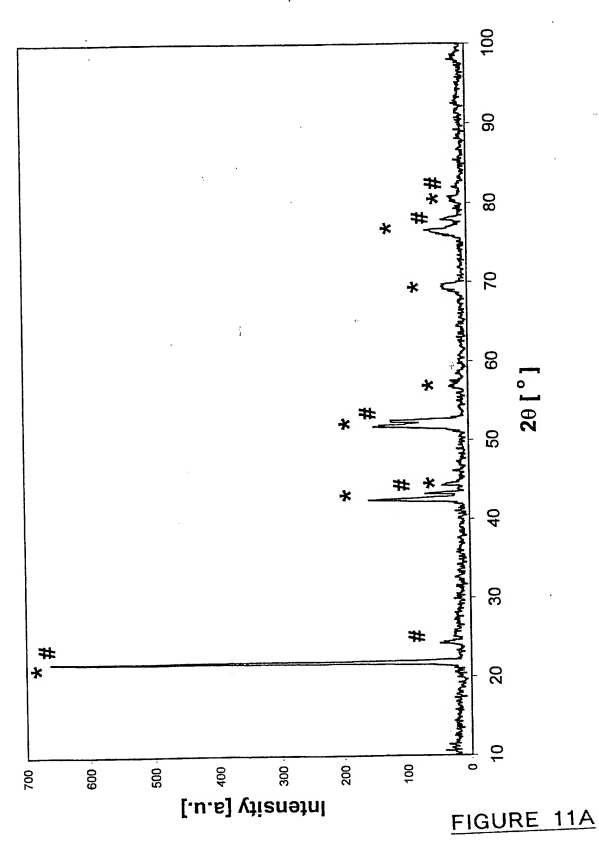
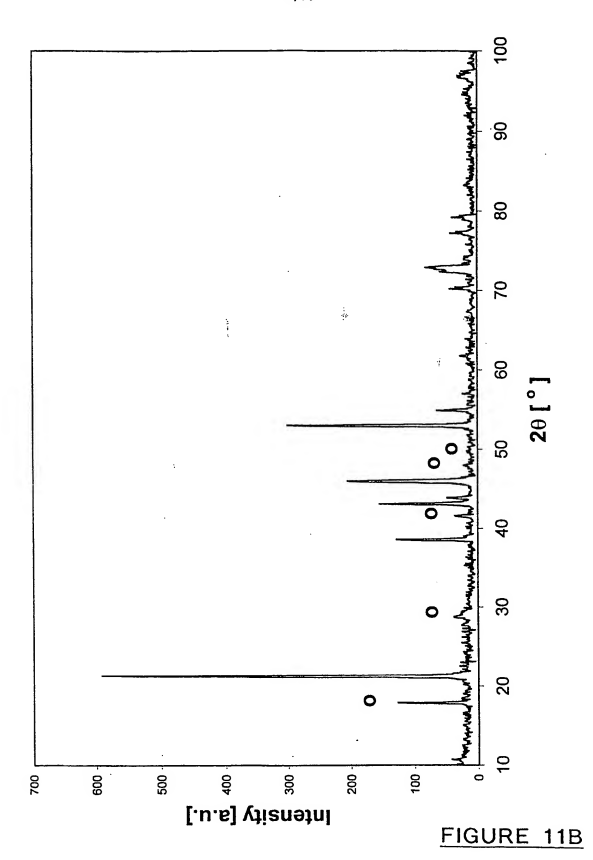
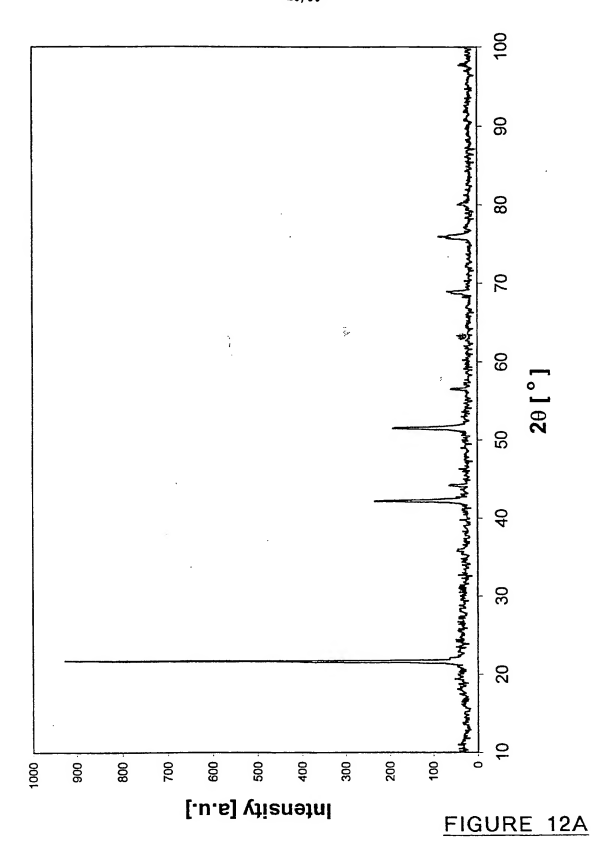
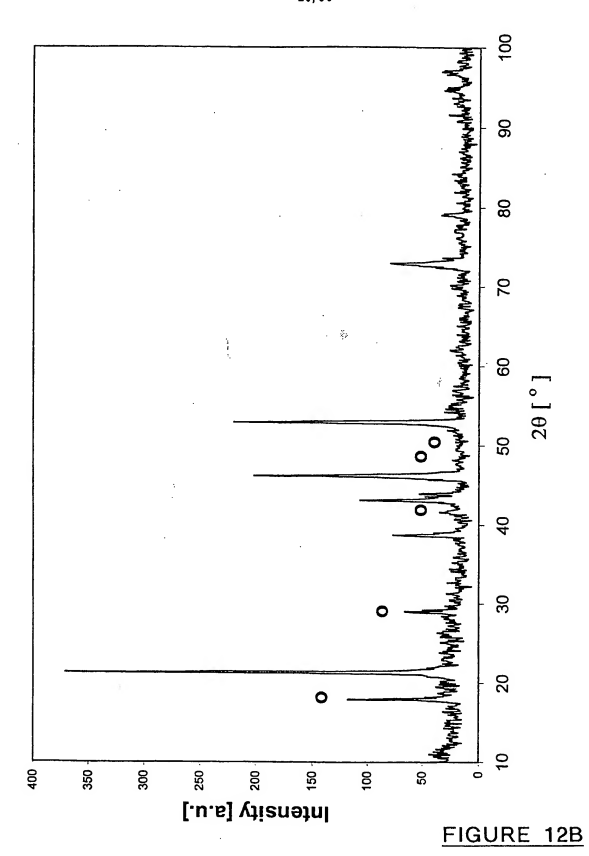


FIGURE 10B









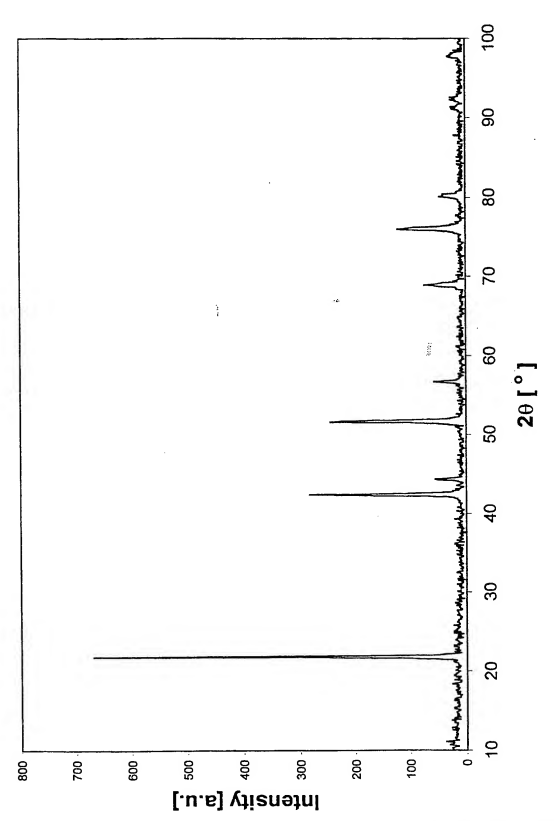
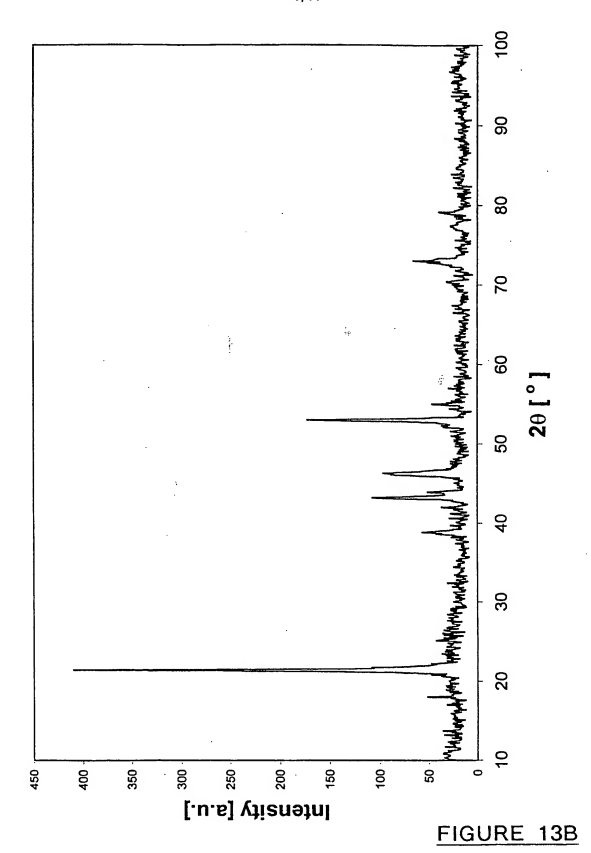


FIGURE 13A



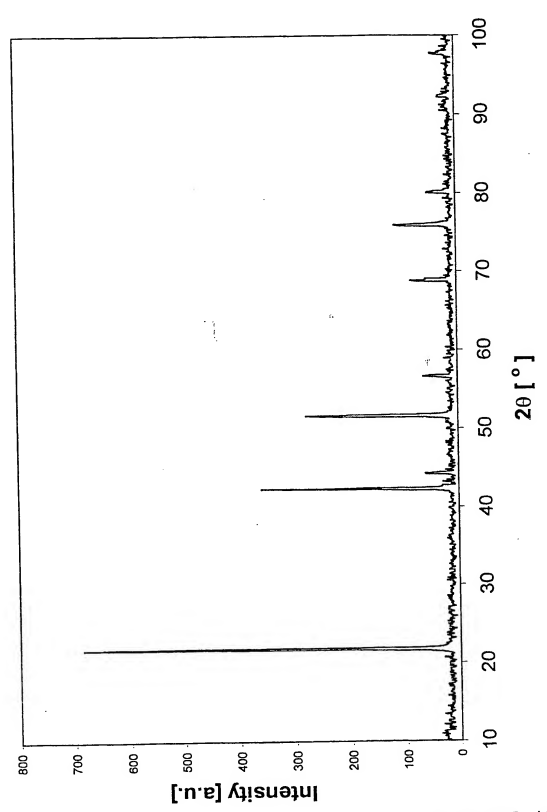


FIGURE 13C

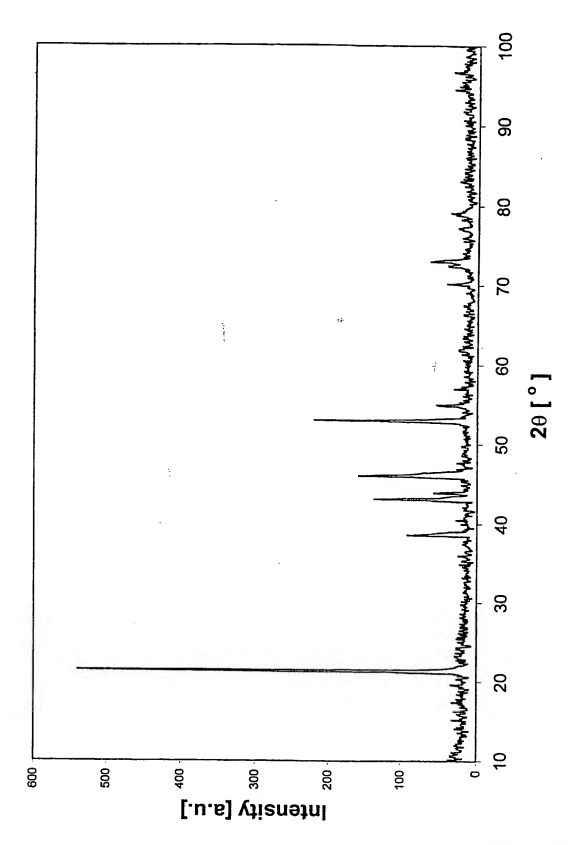


FIGURE 13D

INTERNATIONAL SEARCH REPORT

International application No. PCT/NZ 99/00175

A. (CLASSIFICATION OF SUBJECT MATTER							
Int Cl ⁶ :	C01D 15/02; C01G 45/02; H01M 4/50, 4/58							
According to International Patent Classification (IPC) or to both national classification and IPC								
B. FIELDS SEARCHED								
C01D 15/02,	mentation searched (classification system followed by classify), COIG 45/02, 45/00; H01M 4/50, 4/48,	4/58						
Documentation	searched other than minimum documentation to the exte	nt that such documents are included in the	he fields searched					
Electronic data WPAT, CAS	base consulted during the international search (name of one), JAPIO	data base and, where practicable, search	terms used)					
C.	DOCUMENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.					
х	EP 782206 A (Japan Storage Battery Co) 2 July 1 See whole document	1, 4-5, 54-56						
x	WO 98/02928 A (SAFT) 22 January 1998. See abstract, claims 1&2	1, 4-7, 54-56						
x	Derwent Abstract Accession No. 98-353168, Class L03, JP 10134812 A (Matsushita) 22 May 1998 X See abstract							
x	Further documents are listed in the continuation of Box C	X See patent family ar	nnex					
* Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention cannot be considered novel or cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive								
Date of the ac	tual completion of the international search	Date of mailing of the international search report 1 FEB 2000						
19 January 2	iling address of the ISA/AU	Authorized officer						
AUSTRALIA PO BOX 200 E-mail addres	N PATENT OFFICE , WODEN ACT 2606, AUSTRALIA ss: pct@ipaustralia.gov.au . (02) 6285 3929	JAMES DZIEDZIC Telephone No.: (02) 6283 2495						

INTERNATIONAL SEARCH REPORT

International application No.
PCT/NZ 99/00175

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to							
alegory	Chanon of document, with indication, where appropriate, of the relevant passages	claim No.					
	Derwent Abstract Accession No . 98-039244, Class L03, JP 09293508 A (Sony) 11 November 1997						
X	See abstract	1, 4-7, 54-50					
	Derwent Abstract Accession No. 97-391042 Class X16, JP 09171824 A (Matsushita) 30 June 1997						
х	See abstract	1,4, 54-56					
	Derwent Abstract Accession No. 97-070344, Class L03, JP 08315819 A (Yuasa) 29 November 1996						
X	See abstract	1, 4-5, 54-5					
	Derwent Abstract Accession No. 96-214217 Class L03, JP 08078009 A (Yuasa) 22 March 1996						
X	See abstract	1,4,6, 54-5					
	Derwent Abstract Accession No. 96-214215 Class L03, JP 08078007 A (Yuasa) 22 March 1996						
X	See abstract	1,4,5, 54-5					
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INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No. PCT/NZ 99/00175

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Do	curnent Cited in Search Report			Patent	Family Member		
EP	782206	CN	1156910	JP	9237631	US	5718989
		US	5795558				
wo	98/02928	CA	2228671	EP	858677	FR	2751135
							END OF ANNEX